### Scientific and Technical Information Center

## SEARCH REQUEST FORM

Requester's Full Name: MART Unit: 1624 Phone N	RK BACH Exa  Tumber: 2- 0663	miner #: 59193 Date: 2/22/06 / Serial Number: 10527649
Location (Bldg/Room#): 5 CO1 (N	failbox #): 5C18 Resul	ts Format Preferred (circle): (PAPER) DISK
To ensure an efficient and quality search, ple	ease attach a copy of the cover she	et, claims, and abstract or fill out the following:
Title of Invention:		·
Inventors (please provide full names):		
·		
Earliest Priority Date:		
	ms, acronyms, and registry number	y as possible the subject matter to be searched. Include the rs, and combine with the concept or utility of the invention. ations, authors, etc., if known.
*For Sequence Searches Only* Please include appropriate serial number.	all pertinent information (parent,	child, divisional, or issued patent numbers) along with the
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STAFF USE ONLY	Type of Search	Vendors and cost where applicable
Searcher:	NA Sequence (#)	1286 STNDialog
Searcher Phone #:	AA Sequence (#)	Questel/Orbit Lexis/Nexis
Searcher Location:	Structure (#)	WestlawWWW/Internet
Date Searcher Picked Up:	Bibliographic	in-house sequence systems
Date Completed: 3/3	Litigation	Commercial Oligomer Score/Length Interference SPDI Encode/Transl
Searcher Prep & Review Time: 14	Fulltext	Other (specify)
Online Time: 26	Other	.•

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(FILE 'HOME' ENTERED AT 14:28:20 ON 03 MAR 2006)

FILE 'REGISTRY' ENTERED AT 14:28:31 ON 03 MAR 2006 E CEFDINIR/CN 5

L1 1 S E3
L2 STR 91832-40-5
L3 STR L2
L4 STR L3
L5 7 S L2 OR L3 OR L4
L6 166 S L2 OR L3 OR L4 FUL

L7 SCR 2127 L8 55 SEARCH L7 SUB=L6 FUL

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NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

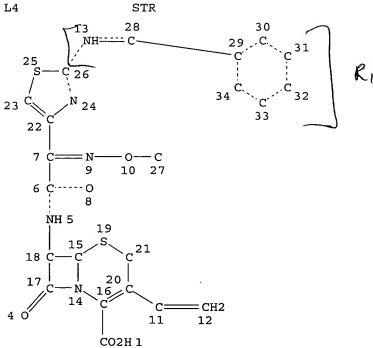
GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 25

STEREO ATTRIBUTES: NONE

NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 27

STEREO ATTRIBUTES: NONE



NODE ATTRIBUTES:

DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED

NUMBER OF NODES IS 32

STEREO ATTRIBUTES: NONE

L6 166 SEA FILE=REGISTRY SSS FUL L2 OR L3 OR L4

L7 SCR 2127

L8 55 SEA FILE=REGISTRY SUB=L6 SSS FUL L7

FULL SUBSET SCREEN SEARCH COMPLETED

55 ANSWERS

SEARCH TIME: 00.00.01

=> fil caplus;s 18 COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION 388.74 388.95

FULL ESTIMATED COST

FILE 'CAPLUS' ENTERED AT 14:32:44 ON 03 MAR 2006
USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.
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FILE COVERS 1907 - 3 Mar 2006 VOL 144 ISS 11 FILE LAST UPDATED: 2 Mar 2006 (20060302/ED)

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http://www.cas.org/infopolicy.html

L9 52 L8

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L9 ANSWER 1 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2006:136151 CAPLUS

DOCUMENT NUMBER:

INVENTOR(S):

144:170821

TITLE:

Preparation of cefixime disodium salt as antibiotic Yu, Anguo; Lin, Guohua; Tang, Chaoyun; Mo, Zhaoming;

Li, Sha

PATENT ASSIGNEE(S):

Peop. Rep. China

Page 4

SOURCE: Faming Zhuanli Shenqing Gongkai Shuomingshu, 6 pp.

CODEN: CNXXEV

DOCUMENT TYPE:

Patent

LANGUAGE:

Chinese

DANGUAGE:

- -

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
CN 1594322	Α	20050316	CN 2004-10040017	20040618
PRIORITY APPLN. INFO.:			CN 2004-10040017	20040618
	<b>a</b> .	<b>-</b> -	11 1 1 1	

AB Cefixime disodium salt, useful as antibiotic, was prepared by treatment of cefixime with NaHCO3. Thus, a mixture of cefixime (507.5 g) and 10% NaHCO3 aqueous solution (1680 g) was stirred for 2 h at rt. Activated carbon (10 g)

was

added and stirring was continued for addnl. 20 min before filtration. The filtrate was treated with ethanol, and the resultant precipitate was collected and dried at 50°C to give crude product, which was recrystd. with ethanol and dried to afford the pure sodium salt in 85.9% yield. This product showed similar antibacterial activity to cefixime and low toxicity.

IT 79350-82-6P

RL: ADV (Adverse effect, including toxicity); IMF (Industrial manufacture); PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of cefixime disodium salt as antibiotics, via neutralization of cefixime with NaHCO3)

RN 79350-82-6 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2-amino-4-thiazolyl)[(carboxymethoxy)imino]acetyl]amino]-3-ethenyl-8-oxo-, disodium salt, [6R-[6 $\alpha$ ,7 $\beta$ (Z)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

●2 Na

L9 ANSWER 2 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2006:79174 CAPLUS

#### Page 5

DOCUMENT NUMBER:

144:170818

TITLE:

Preparation of tertiary amine salts of

2-(2-aminothiazol-4-yl)-2-(acyloxyimino)acetic acid as

intermediates for cefdinir

INVENTOR(S):

Kremminger, Peter; Silberberger, Herbert

PATENT ASSIGNEE(S): .

Sandoz AG, Switz.

SOURCE:

PCT Int. Appl., 18 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

GI

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PA'	TENT	NO.	,		KIN	D .	DATE		1	APPL:	ICAT:	ION I	NO.		Dž	ATE	
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WO	2006	0081	60		A1	:	2006	0126	Ţ	WO 2	005-1	EP79	58		20	0050	721
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							ID,										
							LU,										
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							MC,										
							GN,										
		GM,	KE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	AZ,	BY,
		KG,	KZ,	MD,	RU,	TJ,	TM										
PRIORIT	Y APP	LN.	INFO	. :	•	•			(	GB 2	004-	1637	9	1	A 2	0040	722

Crystalline tertiary amine salts of 2-(2-aminothiazol-4-yl)-2-AB · (acyloxyimino) acetic acid compds. of formula (I) (R1, R2, R3 = independently unsubstituted or substituted alkyl, cycloalkyl or aryl; R4 = acyl) are prepared These salts may be obtained in anhydrous form and are useful in a reaction step with an activating agent in order to produce cefdinir. Thus, 25.0 g syn-2-(2-aminothiazol-4-yl)-2-[[(methylcarbonyl)oxy]imino]acetic acid monohydrate (water content: 8.0%) was suspended in 20 mL acetone at ambient temperature and 5.2 mL tributylamine was added. The mixture was cooled to -10° and stirred at this temperature for 60 and filtered to give, after washing with a small portion of cold acetone and dried in vacuum to give, 32.7 g tributylammonium syn-2-(2-aminothiazol-4-yl)-2-[[(methylcarbonyl)oxy]imino]acetate (water content: 0.1%) (II). II was converted into syn-2-(2-aminothiazol-4-yl)-2-[[(methylcarbonyl)oxy]imino]acetic acid 2-benzothiazolyl thioester by treatment with bis(benzothiazol-2-yl) disulfide and then condensed with 7-amino-3-vinyl-cephem-4-carboxylic acid to give 7-[2-(2-aminothiazol-4-

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Page 6
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IT

y1) -2-[[(methylcarbonyl)oxy]imino]acetamido]-3-vinylcephem-4-carboxylic acid phosphate which was converted into cefdinir by treatment with a mixture of concentrated H2SO4 in MeOH. 663170-79-4P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT

(Reactant or reagent)

(preparation of tertiary amine salts of 2-(2-aminothiazole-4-yl)-2-(acyloxyimino) acetic acid as intermediates for cefdinir)

663170-79-4 CAPLUS RN

5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, CN 7-[[(2Z)-[(acetyloxy)imino](2-amino-4-thiazolyl)acetyl]amino]-3-ethenyl-8oxo-, (6R,7R)-, phosphate (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 127770-93-8 CMF C16 H15 N5 O6 S2

Absolute stereochemistry. Double bond geometry as shown.

CM 2

CRN 7664-38-2 CMF H3 O4 P

IT 874438-71-8P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of tertiary amine salts of 2-(2-aminothiazole-4-yl)-2-(acyloxyimino) acetic acid as intermediates for cefdinir)

874438-71-8 CAPLUS RN

5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, CN 7-[[(2Z)-[(acetyloxy)imino](2-amino-4-thiazolyl)acetyl]amino]-3-ethenyl-8oxo-, (6R,7R)-, mono(4-methylbenzenesulfonate) (9CI) (CA INDEX NAME)

CM 1

CRN 127770-93-8 CMF C16 H15 N5 O6 S2

Absolute stereochemistry.

Double bond geometry as shown.

CM 2

CRN 104-15-4 CMF C7 H8 O3 S

REFERENCE COUNT:

THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 3 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2006:76118 CAPLUS

DOCUMENT NUMBER:

144:170817

TITLE:

Preparation of alkamide solvates of

2-(2-aminothiazol-4-yl)-2-(acyloxyimino)acetic acid as

intermediates for cefdinir

INVENTOR(S):

Kremminger, Peter; Silberberger, Herbert

PATENT ASSIGNEE(S):

Sandoz AG, Switz.

SOURCE:

PCT Int. Appl., 15 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2006008161	A1	20060126	WO 2005-EP7963	20050721
W: AE, AG, AL,	AM, AT	, AU, AZ, BA	, BB, BG, BR, BW, BY,	BZ, CA, CH,

CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NG, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SM, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG, BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM GB 2004-16380 A 20040722

PRIORITY APPLN. INFO.: GI

AB Crystalline N, N-dimethylalkamide solvates of 2-(2-aminothiazole -4-yl)-2(acyloxyimino)acetic acid compds. of formula (I) [R1 = H, (un) substituted alkyl; R4 = acyl] are prepared These compds. may be prepared in an anhydrous form and are useful in a reaction step with an activating agent in order to produce cefdinir. Thus, 15.0 g syn-2-(2-aminothiazol-4y1)-2-[[(methylcarbonyl)oxy]imino]acetic acid dihydrate (H2O content 13.5% ) was dispensed into 54.0 mL N, N-dimethylacetamide at 50° and stirred for 90 min. The crystalline suspension was cooled to 0°, treated with 150 mL CH2Cl2 and the white crystals were filtered, washed three times, each with 30 mL CH2Cl2, and dried over night in vacuum at 30° to give 15,9 g syn-2-(2-aminothiazol-4-yl)-2-[[(methylcarbonyl)oxy]imino]acetic acid N,N-dimethylacetamide solvate (II) (water content 0.4 %). II was converted into syn-2-(2-aminothiazol-4-yl)-2-[[(methylcarbonyl)oxy]imino]acetic acid benzothiazol-2-yl thioester by treatment with bis(benzothiazol-2-yl) disulfide followed by amidation with 7-amino-3-vinylcephem-4-carboxylic acid and acidification with phosphoric acid to give 7-[2-(2-aminothiazol-4-yl)-2-[[(methylcarbonyl)oxy]imino]acet amido]-3-vinylcephem-4-carboxylic acid phosphate (III). Cefdinir was obtained by treatment of III with a mixture of concentrated H2SO4 and MeOH. TT 663170-79-4P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(intermediate; preparation of alkamide solvates of 2-(2-aminothiazol-4-yl)-2-

(acyloxyimino) acetic acid as intermediates for cefdinir)

RN663170-79-4 CAPLUS

5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, CN 7-[[(2Z)-[(acetyloxy)imino](2-amino-4-thiazolyl)acetyl]amino]-3-ethenyl-8oxo-, (6R,7R)-, phosphate (1:1) (9CI) (CA INDEX NAME)

CM1

CRN 127770-93-8 C16 H15 N5 O6 S2 CMF

Absolute stereochemistry.

Double bond geometry as shown.

CM 2

CRN 7664-38-2 CMF H3 O4 P

IT 874438-71-8P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of alkamide solvates of 2-(2-aminothiazol-4-yl)-2-(acyloxyimino)acetic acid as intermediates for cefdinir)

RN 874438-71-8 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,
7-[[(2Z)-[(acetyloxy)imino](2-amino-4-thiazolyl)acetyl]amino]-3-ethenyl-8oxo-, (6R,7R)-, mono(4-methylbenzenesulfonate) (9CI) (CA INDEX NAME)

CM 1

CRN 127770-93-8 CMF C16 H15 N5 O6 S2

Absolute stereochemistry.

Double bond geometry as shown.

CM2

CRN 104-15-4 C7 H8 O3 S CMF

REFERENCE COUNT:

THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

CAPLUS COPYRIGHT 2006 ACS on STN ANSWER 4 OF 52

ACCESSION NUMBER:

2006:54564 CAPLUS

DOCUMENT NUMBER:

144:128794

TITLE:

News salts in the preparation of cephalosporin

antibiotics

INVENTOR(S):

Senthilkumar, Udayampalayam Palanisamy; Lakshmipathi, Venu Sanjeevi; Andrew, Gnanaprakasam; Chandrasekaran, Ramasubbu; Nagender Rao, Dindigala; Om Reddy, Gaddam

PATENT ASSIGNEE(S):

Orchid Chemicals & Pharmaceuticals Limited, India

SOURCE:

PCT Int. Appl., 23 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT	NO.			KIN	D 1	DATE		;	APPL	ICAT	ION 1	NO.		Di	ATE	
					-									-		
WO 2006	00604	40		A2		2006	0119	1	WO 2	005-	IB18	88		2	00501	704
W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	ΒZ,	CA,	CH,
	CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,
	GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KM,	KP,	KR,	KZ,
	LC,	LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	ΜZ,	NA,
	NG,	NI,	NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,
	SL,	SM,	SY,	ТJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	υs,	UZ,	VC,	VN,	YU,
	ZA,	ZM,	ZW													

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RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG, BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM

PRIORITY APPLN. INFO::

IN 2004-CH637 A 20040705
```

The present invention relates to an improved process for the preparation of cephalosporin antibiotics via the formation of intermediate diamine salts of the general form Cp.nM [Cp = cephalosporin antibiotic, such as Cefdinir, Cefoxitin, Cefonicid, etc.; M = ethylenediamine derivative, such as N,N'-diisobutyl-, N,N'-dicyclohexyl-, N,N'-diisopentyl-, N,N'-di(p-anisyl)-, N,N'-dicyclopentyl-, N,N'-di(p-tolyl)-1,2-ethanediamine; n = 0.5 - 2]. Thus, the N,N'-diisobutyl-1,2-ethanediamine salt of Cefonicid (I) was prepd via a reaction of  $7\beta$ -aminocephem II with O-formyl-D-mandeloyl chloride, adjustment of the reaction mixture to pH 5±1, and finally, addition of the diacetate salt of Me2CHCH2NH(CH2)2NHCH2CHMe2.

IT 696592-17-3 717098-27-6

RL: RCT (Reactant); RACT (Reactant or reagent)
(preparation of intermediate salts for the preparation of cephalosporin antibiotics, such as Cefdinir)

RN 696592-17-3 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2Z)-(2-amino-4-thiazolyl)[(triphenylmethoxy)imino]acetyl]amino]-3-ethenyl-8-oxo-, monopotassium salt, (6R,7R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

K

RN 717098-27-6 CAPLUS
CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,
7-[[(2Z)-(2-amino-4-thiazolyl)[(triphenylmethoxy)imino]acetyl]amino]-3-ethenyl-8-oxo-, (6R,7R)-, compd. with N-cyclohexylcyclohexanamine (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 128454-32-0 CMF C33 H27 N5 O5 S2

Absolute stereochemistry. Double bond geometry as shown.

$$Ph_3C$$
 $O$ 
 $R$ 
 $R$ 
 $R$ 
 $R$ 
 $CH_2$ 

CM 2

CRN 101-83-7 CMF C12 H23 N

IT 873441-06-6P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of intermediate salts for the preparation of cephalosporin antibiotics, such as Cefdinir)

RN 873441-06-6 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,
7-[[(2Z)-(2-amino-4-thiazolyl)[(triphenylmethoxy)imino]acetyl]amino]-3ethenyl-8-oxo-, (6R,7R)-, compd. with N,N'-dicyclohexyl-1,2-ethanediamine
(9CI) (CA INDEX NAME)

CM 1

CRN 128454-32-0 CMF C33 H27 N5 O5 S2

Absolute stereochemistry.

Double bond geometry as shown.

CM 2

CRN 4013-98-3 CMF C14 H28 N2

L9 ANSWER 5 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2005:1215707 CAPLUS

DOCUMENT NUMBER:

143:466198

TITLE:

Novel pharmaceutical formulation of cefixime for

enhanced bioavailability

INVENTOR(S): Wagh, Sanjay; Aga, Hidaytulla; Avachat, Makarand; Sen,

Himadri

PATENT ASSIGNEE(S): Lupin Ltd., India

SOURCE: PCT Int. Appl., 18 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATEN	T NO.			KINI	IND DATE APPLICATION NO.						DATE					
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WO ·20	051077	03		A1	:	2005:	1117	1	WO 2	004-	IN128	8		20	040!	510
W	: AE,	AG,	AL,	AM,	ΑT,	AU,	ΑZ,	BA,	BB,	BG,	BR,	BW,	BY,	ΒZ,	CA,	CH,
	CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,
	GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,
	LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NA,	NI,
	NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,
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R	W: BW,	GH,	GM,	KE,	LS,	MW,	MZ,	NA,	SD,	SL,	sz.	TZ.	ŪĠ,	ZM.	ZW,	AM,
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AB A chewable tablet comprises cefixime having a mean particle size 20-120  $\,$   $\,$   $\,$  µm, wherein the composition demonstrates bioequivalence to a suspension of cefixime trihydrate. The process of preparing the chewable tablet comprises the steps of optionally micronizing cefixime such that the mean particle size of the cefixime particles is 20-120  $\,$  µm, blending with other excipients, roll compaction, milling to form granules, blending to form a secondary blend and compression of the secondary blend to form tablets.

IT 125110-14-7, Cefixime trihydrate

RL: PKT (Pharmacokinetics); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(chewable tablets containing cefixime with enhanced bioavailability)

RN 125110-14-7 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,

7-[[(2Z)-(2-amino-4-thiazolyl)[(carboxymethoxy)imino]acetyl]amino]-3-ethenyl-8-oxo-, trihydrate, (6R,7R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

●3 H<sub>2</sub>O

REFERENCE COUNT: 7 THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 6 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2005:823155 CAPLUS

DOCUMENT NUMBER:

143:235396

TITLE:

Synergistic antibacterial formulation containing

cefixime trihydrate, cloxacillin sodium and

Lactobacillus sporogenes spores

INVENTOR(S):

Khandelwal, Sanjeev

PATENT ASSIGNEE(S):

India

SOURCE:

U.S. Pat. Appl. Publ., 25 pp.

CODEN: USXXCO

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PAT	ENT 1	. 01			KIN	)	DATE			APP:	LICAT	ION I	NO.		D	ATE	
						-						-			-		
US :	20051	1810	51		A1		2005	0818		US :	2004-	1311	0		2	0041	215
EP	15661	176			A1		2005	0824		EP :	2005-	2508	79		2	0050	216
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR	, IT,	LI,	LU,	NL,	ŞΕ,	MC,	PT,
		ΙE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL	, TR,	BG,	CZ,	EE,	HU,	PL,	SK,
		BA,	HR,	IS,	YU												
PRIORITY	APPI	LN.	INFO	. :						IN :	2004-	MU17	8	1	A 2	0040	216
										IN :	2004-	MU25	8	1	A 2	0040	303

As synergistic antibacterial formulation for oral delivery of cefixime trihydrate, cloxacillin sodium in an extended release form and an immediate release form, and Lactobacillus sporogenes spores is provided. For example, sustained-release granules were prepared by wet granulation of cloxacillin sodium 50.0 kg and hydroxypropyl Me cellulose (HPMC; average viscosity 4000 cps) 6.0 kg, using a binder comprising HPMC (average viscosity 50 cps) 800g dissolved in a mixture of dichloromethane 8.0 kg and iso-Pr alc. 12.0 kg. The core was prepared by blending cloxacillin sodium sustained-release granules obtained with a mixture of cloxacillin sodium particle 7.6 kg, cefixime trihydrate particles 11.2 kg, L. sporogenes spores 750 g, sodium starch glycollate 1.0 kg, colloidal silicon dioxide

0.3 kg, sodium lauryl sulfate 1.0 kg and talc 1.0 kg was prepared Magnesium stearate 1.0 kg was added and further blendded, resulting in the lubricated core mass. This core mass was then compressed into cores of average weight of 806.2 mg <plus/minus>3%. The core obtained were pan coated with a film coating composition containing Et cellulose 0.8 kg, hydroxypropyl cellulose 0.8 kg, iso-Pr alc. 12 kg, methylene chloride 22 kg, di-Et phthalate 0.01 kg and titanium dioxide 0.15 kg in a stainless steel container and stirred for five minutes using overhead stirrer until a smooth slurry was obtained. The coated tablets were polished with talc. The film-coated tablet (average weight 820 mg <plus/minus>3%) contained (i) cloxacillin sodium equivalent to 250 mg cloxacillin sustained release, (ii) cloxacillin sodium equivalent to 250 mg cloxacillin immediate release, (III) cefixime trihydrate equivalent to 100 mg cefixime immediate release, and (IV) L. sporogenes 45 million spores.

IT 125110-14-7, Cefixime Trihydrate

RL: ADV (Adverse effect, including toxicity); PAC (Pharmacological activity); PKT (Pharmacokinetics); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(synergistic antibacterial formulation containing cefixime trihydrate, cloxacillin sodium and Lactobacillus sporogenes spores)

RN 125110-14-7 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,

7-[[(2Z)-(2-amino-4-thiazolyl)[(carboxymethoxy)imino]acetyl]amino]-3-ethenyl-8-oxo-, trihydrate, (6R,7R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

$$HO_2C$$
 $O$ 
 $R$ 
 $R$ 
 $R$ 
 $R$ 
 $CH_2$ 

●3 H<sub>2</sub>O

L9 ANSWER 7 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2005:315131 CAPLUS

DOCUMENT NUMBER: 142:336175

TITLE: An improved process for the preparation of cefixime

trihydrate

INVENTOR(S): Sharma, Anil Kumar; Raj, Baldev; Sethi, Madhuresh

Kumar; Das, Debashis

PATENT ASSIGNEE(S): J K Drugs & Pharmaceuticals Ltd., India

SOURCE: Port. Pat. Appl., 27 pp.

CODEN: PTXXB9

Page 17

DOCUMENT TYPE:

Patent

LANGUAGE:

Portuguese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
				-	
PT 102293	Α	20000229	PT 1999-102293.		19990426
PT 102293	В	20010531			
IN 185070	A	20001104	IN 1999-BO75		19990129
PRIORITY APPLN. INFO.:			IN 1999-BO75	Α	19990129
OTHER SOURCE(S):	CASREA	CT 142:33617	5; MARPAT 142:336175		•
GI	•				

- STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY -AVAILABLE VIA OFFLINE PRINT \*
- An improved process for the preparation of cefixime trihydrate (I.3H2O) AB comprises: (a) hydrolysis of the 3-acetoxymethyl group of 7-(substituted amino) cephalosporanic acid [II; R = H, CO(CH2)3CH(NH2)CO2H] with an alkali carbonate; (b) protective acylation of the 7-amino group with an organic acid chloride; (c) esterification of the 4-carboxy group; (d) bromination of the 3-hydroxymethyl group with PBr3; (e) Wittig reaction with HCHO in the presence of PPh3 to give a 3-vinyl compound III; (f) cleavage of the phenylacetyl group from the 7-amino group with the PPh3/Cl2/pyridine/IBA complex; (g) acylation of the resulting 7-amino group with 4-chloro-2-[{ (methoxycarbonyl) methoxy}imino]-3-oxobutyric acid; (h) cyclization of the acylated cephem IV (R1 = CHPh2, CH2C6H4OMe) with thiourea to give protected I; and (i) removal of the protective group.
- 125110-14-7P, Cefixime trihydrate IT RL: SPN (Synthetic preparation); PREP (Preparation) (improved process for preparation of cefixime trihydrate)
- 125110-14-7 CAPLUS RN
- CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2Z)-(2-amino-4-thiazolyl)[(carboxymethoxy)imino]acetyl]amino]-3ethenyl-8-oxo-, trihydrate, (6R,7R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

#### ●3 H<sub>2</sub>O

L9 ANSWER 8 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2004:1038759 CAPLUS

DOCUMENT NUMBER: 142:232241

TITLE: Spectrophotometric determination of some

cephalosporins in biological fluids using ferric-phenanthroline and tetrazolium blue

AUTHOR(S): Abdel-Razeq, Sawsan A.

CORPORATE SOURCE: Pharmaceutical Chemistry Department, Pharmacy College,

Al-Azhar University, Cairo, Egypt

SOURCE: Bulletin of the Faculty of Pharmacy (Cairo University)

(2002), 40(1), 155-166

CODEN: BFPHA8; ISSN: 1110-0931

PUBLISHER: Cairo University, Faculty of Pharmacy

DOCUMENT TYPE: Journal LANGUAGE: English

AB Two sensitive spectrophotometric procedures are presented for the determination of

three cephalosporins; cefixime trihydrate (I), cefoperazone sodium (II) and cefotaxime sodium (III). The first procedure is based on the reduction of ferric into ferrous in presence of o-phenanthroline by the mentioned drugs to form a highly stable orange-red ferroin chelate [Fe-(Phen)3]2+, measured at 513 nm. The second procedure is also based on the reduction of tetrazolium blue (TZB) in alkaline medium by the above cephalosporins leading to the formation of highly colored purple formazan measured at 526 nm. Beer's law is obeyed in the ranges of 0.4 - 2.4 and 4-20  $\mu g$  ml-1 for I, 0.8 - 3.6 and 4 - 24  $\mu g$  ml-1 for II or 0.4 - 2.4 and 4 - 16  $\mu g$  ml-1 for III by Ferric- phen and TZB procedures, resp. The optimum assay conditions and their applicability to the determination of the cited drugs in pharmaceutical formulations are described. The recoveries of the drugs are 90.7-96.0% from urine and 71.7 - 78.5% from serum.

IT 125110-14-7, Cefixime trihydrate

RL: BSU (Biological study, unclassified); BIOL (Biological study) (spectrophotometry methods using ferric-phenanthroline and tetrazolium blue are effective and sensitive in determining cephalosporin cefixime trihydrate in biol. fluid)

RN 125110-14-7 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,

7-[[(2Z)-(2-amino-4-thiazolyl)[(carboxymethoxy)imino]acetyl]amino]-3-ethenyl-8-oxo-, trihydrate, (6R,7R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

●3 H<sub>2</sub>O

REFERENCE COUNT:

THERE ARE 18 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 9 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

18

ACCESSION NUMBER:

2004:546513 CAPLUS

DOCUMENT NUMBER:

141:88964

TITLE:

Process for preparing crystalline cefdinir salts Pozzi, Giovanni; Martin Gomez, Patricio; Alpegiani,

INVENTOR(S):

Marco; Cabri, Walter

PATENT ASSIGNEE(S):

Antibioticos S.p.A., Italy PCT Int. Appl., 14 pp.

SOURCE:

CODEN: PIXXD2

C

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA'	rent :	NO.			KIN	<b>)</b>	DATE		1	APPL	ICAT:	ION I	. O <i>v</i>		D	ATE		-
 ₩O	2004	 0568			Δ1		 2004:	0708	,	 WO 21	003-1	 ED13	 524		. 20	0031	201	
***		AE,																
		CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,	
		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	ΚP,	KR,	KZ,	LC,	
		LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	ΝÍ,	NO,	
		NZ,	OM,	PG,	ΡĤ,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,	ТJ,	
		TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UΖ,	VC,	VN,	YU,	ZA,	ZM,	ZW		
	RW:	BW,	GH,	GM,	KΕ,	LS,	MW,	ΜZ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	AZ,	
		BY,	KG,	KZ,	MD,	RU,	ТJ,	TM,	ΑT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	
		ES,	FI,	FR,	GB,	GR,	HU,	ΙE,	IT,	LU,	MC,	ΝL,	·PT,	RO,	SE,	SI,	SK,	
		TR,	BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	ΝE,	SN,	TD,	TG
EP	1572	699			A1		2005	0914		EP 2	003-	7891	09		2	0031	201	
	R:	AT,															PT,	
		IE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	ΑL,	TR,	BG,	CZ,	EE,	HU,	SK		

PRIORITY APPLN. INFO.:

IT 2002-MI2724 WO 2003-EP13524 A 20021220 W 20031201

OTHER SOURCE(S):

MARPAT 141:88964

GI

AB Cefdinir salts, such as I.nH3PO4 [R1, R2 = H; n = 1 - 3 (II)], the hydrates and solvates thereof, were prepared from cefdinir intermediates, I (R1 = benzhydryl, trityl, p-methoxybenzyl; R2 = benzhydryl, tert-Bu, p-methoxybenzyl), or crude cefdinir I (R1, R2 = H) by the treatment with phosphoric acid. Thus, I (R1 = CPh3, R2 = H) was dissolved in 85% phosphoric acid and acetonitrile, and reaction mixture was heated at 45°C for 2 h, to afford cefdinir phosphate. The use of II for the preparation and purification of cefdinir is also disclosed.

IT 717098-27-6

RL: RCT (Reactant); RACT (Reactant or reagent)
(preparation and use of cefdinir phosphates for preparing and purification

οf

cefdinir)

RN 717098-27-6 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2Z)-(2-amino-4-thiazolyl)[(triphenylmethoxy)imino]acetyl]amino]-3-ethenyl-8-oxo-, (6R,7R)-, compd. with N-cyclohexylcyclohexanamine (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 128454-32-0

CMF C33 H27 N5 O5 S2

Absolute stereochemistry. Double bond geometry as shown.

CM

CRN 101-83-7 C12 H23 N CMF

NH

REFERENCE COUNT:

THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 10 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2004:453223 CAPLUS

DOCUMENT NUMBER: 141:6966

TITLE: Process for preparing cefdinir and its amorphous

hydrate

Deshpande, Pandurang Balwant; Khadangale, Bhausaheb INVENTOR(S):

Pandharinath; Ramasubbu, Chandrasekaran

PATENT ASSIGNEE(S): Orchid Chemicals & Pharmaceuticals Ltd., India

SOURCE: PCT Int. Appl., 26 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent English LANGUAGE:

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PATENT (	NO.		KINI	) I	DATE			APPL:	ICAT:	ION I	NO.		D	ATE	
				-											
WO 2004	046154		A1	•	2004	0603	1	WO 2	003-	IB50	32.	•	20	0031	110
W:	AE, AG,	AL,	AM,	AT,	AU,	ΑZ,	BA,	BB,	BG,	BR,	BW,	BY,	ΒZ,	CA,	CH,
	CN, CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,
	GH, GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	ΚP,	KR,	ΚZ,	LC,	LK,
	LR, LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NI,	NO,	NZ,
	OM, PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,	ТJ,	TM,
	TN, TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	YU,	ZA,	ZM,	zw		
RW:	GH, GM,	ΚE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	AZ,	ΒY,
	KG, KZ,	MD,	RU,	TJ,	TM,	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,
	FI, FR,	GB,	GR,	HU,	ΙE,	IT,	LU,	MC,	NL,	PT,	RO,	SE,	ŚΙ,	SK,	TR,
	BF, BJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	ΝE,	SN,	TD,	TG
PRIORITY APP	LN. INFO	. :						IN 2	002-1	MA84	8	3	A 2	0021	115
								IN 2	003-1	MA15	2	1	A 2	0030	226
OTHER SOURCE	(S):		CASI	REAC'	T 14	1:69	66;	MARP	AT 1	41:6	966				

AB The present invention discloses a process for preparing cefdinir [I; R1 = H; R2 = CO2H (II)] and its monohydrate via condensing 7-amino-3-cephem-4-carboxylic acid with III (X = ester, thioester, halo, etc.) in the presence of a tertiary amine and an organic solvent, followed by treatment with a base to produce I [R1 = C(Ph)3; R2 = carboxylate ion (IV)], and hydrolyzing IV, using an acid in the presence of a solvent, to produce II. Thus, reaction between III (X = OH) and 2-mercapto-5-phenyl-1,3,4-oxadiazole yielded 2-mercapto-5-phenyl-1,3,4-oxadiazolyl-(Z)-(2-aminothiazol-4-yl)-2-(trityloxyimino) acetate, which, on condensation with 7-amino-3-vinyl-3-cephem-4-carboxylic acid and subsequent hydrolysis, afforded II.

IT 696592-17-3P

RL: IMF (Industrial manufacture); RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (preparation of cefdinir and its amorphous hydrate)

RN 696592-17-3 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2Z)-(2-amino-4-thiazolyl)[(triphenylmethoxy)imino]acetyl]amino]-3-ethenyl-8-oxo-, monopotassium salt, (6R,7R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

K

L9 ANSWER 11 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2004:355098 CAPLUS

#### Page 23

DOCUMENT NUMBER:

140:375021

TITLE:

Intermediate cefdinir salts

INVENTOR(S):

Pozzi, Giovanni; Martin Gomez; Patricio; Alpegiani,

Marco; Cabri, Walter

PATENT ASSIGNEE(S):

Antibioticos S.P.A., Italy PCT Int. Appl., 15 pp.

SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

KIND DATE APPLICATION NO. DATE PATENT NO. ----\_\_\_\_\_\_\_\_\_\_\_\_ 20030926 WO 2004035800 A2 20040429 WO 2003-EP10718 WO 2004035800 20040826 Α3 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
2500791
AA 20040429
CA 2003-2500791
20030926 20040429 CA 2003-2500791 20030926 CA 2500791 AΑ 20050629 EP 2003-788921 · 20030926 EP 1546155 A2 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK JP 2004-544046 JP 2006501305 T220060112 20030926 IT 2002-MI2076 A 20021001 PRIORITY APPLN. INFO .: WO 2003-EP10718 W 20030926

OTHER SOURCE(S):

MARPAT 140:375021

GΙ

$$H_2N$$
 $S$ 
 $OH$ 
 $H_2N$ 
 $S$ 
 $OH$ 
 $N$ 
 $S$ 
 $CO_2H$ 
 $II$ 

AB Disclosed are salts of the general formula (I) wherein R1 is H or an amino-protecting group, R2 is and OH-protecting group, and B is NH3 or an organic base, and a process for the preparation thereof. These salts are useful

intermediates for the preparation of cefdinir (II).

IT 682357-23-9P 683226-97-3P

RL: PRP (Properties); PUR (Purification or recovery); SPN (Synthetic preparation); PREP (Preparation)

(intermediate cefdinir salts)

RN 682357-23-9 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,
7-[[(2-amino-4-thiazolyl)[(triphenylmethoxy)imino]acetyl]amino]-3-ethenyl8-oxo-, (6R,7R)-, compd. with N-cyclohexylcyclohexanamine (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 682357-22-8 CMF C33 H27 N5 O5 S2

Absolute stereochemistry.

Double bond geometry unknown.

CM 2

CRN 101-83-7 CMF C12 H23 N

RN 683226-97-3 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,
7-[[(2Z)-(2-amino-4-thiazolyl)[(triphenylmethoxy)imino]acetyl]amino]-3ethenyl-8-oxo-, (6R,7R)-, compd. with (αR)-αmethylbenzenemethanamine (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 128454-32-0 C33 H27 N5 O5 S2 CMF

Absolute stereochemistry. Double bond geometry as shown.

CM

CRN 3886-69-9 CMF C8 H11 N

Absolute stereochemistry. Rotation (+).

CAPLUS COPYRIGHT 2006 ACS on STN ANSWER 12 OF 52

ACCESSION NUMBER:

2004:353145 CAPLUS

DOCUMENT NUMBER:

140:357115

TITLE:

Process for the preparation of Cefixime

INVENTOR(S):

Deshpande, Pandurang Balwant; Das, Gautam Kumar;

Deshpande, Pramod Narayan; Chandrasekaran, Ramasubbu;

Ramar, Padmanabhan; Jeyakumar, John Muthiah Raja

Orchid Chemicals and Pharmaceuticals Limited, India

PATENT ASSIGNEE(S): SOURCE:

U.S. Pat. Appl. Publ., 3 pp.

CODEN: USXXCO

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO.	DATE
US 2004082560	A1 20040429	US 2002-310177	20021205
US 6800755	B2 20041005		
WO 2004037832	A1 20040506	WO 2002-IB5313	20021210
W: AE, AG, AL,	AM, AT, AU, AZ,	BA, BB, BG, BR, BY, BZ,	CA, CH, CN,
CO, CR, CU,	CZ, DE, DK, DM,	DZ, EC, EE, ES, FI, GB,	GD, GE, GH,
GM, HR, HU,	ID, IL, IN, IS,	JP, KE, KG, KP, KR, KZ,	LC, LK, LR,

LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG AU 2002348784 Α1 20040513 AU 2002-348784 20021210 PRIORITY APPLN. INFO.: IN 2002-MA785 20021024 WO 2002-IB5313 W 20021210 OTHER SOURCE(S): CASREACT 140:357115; MARPAT 140:357115

GΙ

AB This invention provides an improved process for the preparation of Cefixime I (R = H) with an improved quality having/possessing better color and solubility Thus, ester I (R = Me) was treated with sodium bicarbonate in water and Et acetate followed by a 15% NaOH solution and subsequent adjustment of the pH of the soln to 4.8-5.0 with 19% aqueous HCl solution and further pH adjustment of

the aqueous layer to 2.45-2.55 with 8-10% HCl to give the desired Cefixime in pure form.

IT125110-14-7P, Cefixime trihydrate

> RL: IMF (Industrial manufacture); SPN (Synthetic preparation); PREP (Preparation)

(process for the preparation of Cefixime)

RN 125110-14-7 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2Z)-(2-amino-4-thiazolyl)[(carboxymethoxy)imino]acetyl]amino]-3ethenyl-8-oxo-, trihydrate, (6R,7R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

#### ●3 H<sub>2</sub>O

REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 13 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2004:162698 CAPLUS

DOCUMENT NUMBER:

140:217437

TITLE: INVENTOR(S): Process for the preparation of cefdinir intermediate

Kremminger, Peter; Wolf, Siegfried; Ludescher,

Johannes

PATENT ASSIGNEE(S):

SOURCE:

Sandoz G.m.b.H., Austria

PCT Int. Appl., 37 pp. CODEN: PIXXD2

DOCUMENT TYPE:

LANGUAGE:

Patent English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO.	DATE
		0226. WO 2003-EP8944	20030812
W: AE, AG, AI	, AM, AT, AU,	AZ, BA, BB, BG, BR, BY,	BZ, CA, CH, CN,
		DM, DZ, EC, EE, ES, FI,	
		JP, KE, KG, KP, KR, KZ,	
		NI, NO, NZ, OM, PG, PH,	
		TM, TN, TR, TT, UA, US,	
ZA, ZW		, , , , , , , , , , , , , , , , , , , ,	
	, KG, KZ, MD,	RU, TJ, TM, AT, BE, BG,	CH, CY, CZ, DE,
		GR, HU, IE, IT, LU, MC,	
SI, SK, TH			
		0303 AU 2003-255424	20030812
EP 1554289		0720 EP 2003-787771	
R: AT. BE. CH		FR, GB, GR, IT, LI, LU,	
		MK, CY, AL, TR, BG, CZ,	
JP 2006500356		0105 JP 2004-528469	
		0202 US 2005-524397	
PRIORITY APPLN. INFO.:	i i	AT 2002-1223	A 20020813
THEORET THE PART OF THE PART O	•	AT 2002-1588	
		WO 2003-EP8944	

OTHER SOURCE(S):

MARPAT 140:217437

GT

AB A process is claimed for the synthesis of 7-[2-(2-aminothiazol-4-yl)-2-(methylcarbonyloxyimino)acetamido]-3-vinyl-cephem-4-carboxylic acid (I), in the form of a crystalline salt, such as I.HX [X = Cl-, HSO4-,RYO3-, H2NSO3-, 1/2(SO4)2-; R = alkyl, aryl; Y = S, P], and their use in the preparation of pure cefdinir. Thus, a reactive derivative of syn-2-(2-aminothiazol-4-yl)2-(methylcarbonyloxyimino)-acetic acid, e.g., syn-2-(2-aminothiazol-4-yl)2-(methylcarbonyloxyimino)-acetic acid mercapto-benzothiazolyl ester is reacted with 7-amino-3-vinyl-3-cephem-4-carboxylic acid in silylated form to obtain I, in which the carboxylic acid is optionally silylated. In another aspect, the present invention relates to salt of I, optionally in crystalline form, wherein the salt is selected from the group consisting of phosphate, hydrogen phosphate, mesylate, tosylate, sulfate, hydrogen sulfate and sulfamate.

IT 663170-77-2P 663170-78-3P 663170-79-4P

RL: IMF (Industrial manufacture); PRP (Properties); SPN (Synthetic preparation); PREP (Preparation)

(preparation and X-ray diffraction measurements of intermediates in the production of cefdinir)

RN 663170-77-2 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2Z)-[(acetyloxy)imino](2-amino-4-thiazolyl)acetyl]amino]-3-ethenyl-8-oxo-, (6R,7R)-, sulfate (2:1) (9CI) (CA INDEX NAME)

CM 1

CRN 127770-93-8 CMF C16 H15 N5 O6 S2

Absolute stereochemistry.

Double bond geometry as shown.

CM 2

CRN 7664-93-9 CMF H2 O4 S

RN 663170-78-3 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,
7-[[(2Z)-[(acetyloxy)imino](2-amino-4-thiazolyl)acetyl]amino]-3-ethenyl-8oxo-, (6R,7R)-, sulfate (1:1) (9CI) (CA INDEX NAME)

٤.

CM 1

CRN 127770-93-8 CMF C16 H15 N5 O6 S2

Absolute stereochemistry. Double bond geometry as shown.

CM 2

CRN 7664-93-9 CMF H2 O4 S

RN 663170-79-4 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2Z)-[(acetyloxy)imino](2-amino-4-thiazolyl)acetyl]amino]-3-ethenyl-8-oxo-, (6R,7R)-, phosphate (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 127770-93-8 CMF C16 H15 N5 O6 S2

Absolute stereochemistry. Double bond geometry as shown.

CM 2

CRN 7664-38-2 CMF H3 O4 P

IT 443874-49-5P 663170-80-7P 663170-81-8P

663170-82-9P

RL: IMF (Industrial manufacture); SPN (Synthetic preparation); PREP

(Preparation)

(process and intermediates in the production of cefdinir)

RN 443874-49-5 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2Z)-[(acetyloxy)imino](2-amino-4-thiazolyl)acetyl]amino]-3-ethenyl-8-oxo-, monohydrochloride, (6R,7R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

● HCl

RN 663170-80-7 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2Z)-[(acetyloxy)imino](2-amino-4-thiazolyl)acetyl]amino]-3-ethenyl-8-oxo-, (6R,7R)-, phosphonate (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 127770-93-8 CMF C16 H15 N5 O6 S2

Absolute stereochemistry.

Double bond geometry as shown.

CM 2

CRN 13598-36-2 CMF H3 O3 P

ONE OR MORE TAUTOMERIC DOUBLE BONDS NOT DISPLAYED IN THE STRUCTURE

663170-81-8 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,
7-[[(2Z)-[(acetyloxy)imino](2-amino-4-thiazolyl)acetyl]amino]-3-ethenyl-8oxo-, (6R,7R)-, monosulfamate (9CI) (CA INDEX NAME)

CM 1

CRN 127770-93-8 CMF C16 H15 N5 O6 S2

Absolute stereochemistry.

Double bond geometry as shown.

CM 2

CRN 5329-14-6 CMF H3 N O3 S

RN 663170-82-9 CAPLUS
CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,
7-[[(2Z)-[(acetyloxy)imino](2-amino-4-thiazolyl)acetyl]amino]-3-ethenyl-8oxo-, (6R,7R)-, monobenzenesulfonate (9CI) (CA INDEX NAME)

Page 33

CM 1

CRN 127770-93-8 CMF C16 H15 N5 O6 S2

Absolute stereochemistry.

Double bond geometry as shown.

CM 2

CRN 98-11-3 CMF C6 H6 O3 S

REFERENCE COUNT:

THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 14 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

8

ACCESSION NUMBER:

2004:18736 CAPLUS

DOCUMENT NUMBER:

140:65237

TITLE:

Extended-release drug delivery systems of cefixime

trihydrate

INVENTOR(S):

Khandelwal, Sanjeev; Omray, Pratibha

PATENT ASSIGNEE(S):

India

SOURCE:

U.S. Pat. Appl. Publ., 10 pp.

CODEN: USXXCO

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 2004005361	A1	20040108	US 2003-456690	20030606
PRIORITY APPLN. INFO.:		•	IN 2002-MU506 A	20020706

AB An extended-release oral drug delivery system comprises as active ingredient cefixime trihydrate (I) in combination with a hydrophilic matrix system, and optionally containing addnl. pharmaceutically acceptable constituents, wherein at least 20 % up to but not more than 40 % of I is released from the matrix within 1 h from oral administration and the remainder of the pharmaceutical agent is released at a sustained rate. Granules were prepared from a mixture containing I 30.36, lactose 4.27, starch 2.99, genistein 0.05, and PVP 0.7 kg, then mixed with HPMC 6.25, Et cellulose 0.5, talc 0.35, and Mg stearate 0.35 kg. The lubricated granules were compressed to give tablets and sprayed with a homogeneous solution containing methylene chloride, isopropanol, HPMC, Et cellulose, titania,

plasticizers, and ethanol, to give coated tablets (containing 200 mg I/tablet).

IT 125110-14-7, Cefixime trihydrate

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (extended-release tablets containing cefixime trihydrate in hydrophilic matrix)

RN 125110-14-7 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2Z)-(2-amino-4-thiazolyl)[(carboxymethoxy)imino]acetyl]amino]-3-ethenyl-8-oxo-, trihydrate, (6R,7R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

●3 H<sub>2</sub>O

L9 ANSWER 15 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2003:376868 CAPLUS

DOCUMENT NUMBER: 138:385207

TITLE: A process for the preparation of cefixime via

alkylsulfonate or arylsulfonate salts

INVENTOR(S): Cabri, Walter; Alpegiani, Marco; Pozzi, Giovanni;

Martin Gomez, Patricio; Oliva, Francesco

PATENT ASSIGNEE(S): Antibioticos S.P.A., Italy

SOURCE: PCT Int. Appl., 21 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

```
APPLICATION NO.
                                                                     DATE
     PATENT NO.
                         KIND
                                DATE .
                         _ _ _ _
    WO 2003040148
                          Α1
                                20030515
                                             WO 2002-EP11405
                                                                     20021011
             AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
             CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
             GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
             LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH,
             PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ,
             UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD,
             RU, TJ, TM
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, BG,
             CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,
             PT, SE, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR,
             NE, SN; TD, TG
     EP 1442044
                                 20040804
                                             EP 2002-782888
                                                                     20021011
                          A1
             AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, SK
     JP 2005508387
                          T2
                                20050331
                                             JP 2003-542194
                                                                     20021011
                                             US 2004-494700
     US 2005032771
                          Α1
                                20050210
                                                                     20040927
PRIORITY APPLN. INFO.:
                                             IT 2001-MI2364
                                                                  Α
                                                                    20011109
                                             WO 2002-EP11405
                                                                 W
                                                                     20021011
OTHER SOURCE(S):
                         CASREACT 138:385207; MARPAT 138:385207
GΙ
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\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

```
Cefixime (I) is prepared in high yield and selectivity by: (A) the amidation
     of a 7-amino-3-vinyl-3-cephem-4-carboxylic acid derivative [II; R1 = H, silyl;
    R2 = H, silyl, tert-Bu, 4-methoxybenzyl, 3,4-dimethoxybenzyl, benzhydryl,
    bis(p-methoxyphenyl)methyl] with a 2-(aminothiazol-4-yl)-2-
     (carboxymethoxyimino) acetic acid derivative [III; R3 = H, trityl,
     tert-butoxycarbonyl; 4-methoxybenzyloxycarbonyl; R4 = tert-Bu,
     p-methoxybenzyl, 3,4-dimethoxybenzyl, benzhydryl,bis(4-
     methoxyphenyl)methyl, trityl; Z = carboxy-activating group] to give a
     7-[2-(aminothiazol-4-yl)-2-(carboxymethoxyimino)acetamido]-3-vinyl-3-
     cephem-4-carboxylic acid derivative (IV); (B) directly reacting IV with a
     sulfonic acid RSO3H [R = C1-6 (un)branched chain, Ph, naphthyl] to give
     the cefixime salt (I·RSO3H·nH2O; n = 0-5); and (C)
     converting I·RSO3H·nH2O into I.
     524925-12-0P, Cefixime methanesulfonate monohydrate
     524925-13-1P, Cefixime methanesulfonate
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (in a process for the preparation of cefixime via alkylsulfonate or
        arylsulfonate salts)
RN
     524925-12-0 CAPLUS
     5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,
     7-[[(2Z)-(2-amino-4-thiazolyl)[(carboxymethoxy)imino]acetyl]amino]-3-
     ethenyl-8-oxo-, (6R,7R)-, mononmethanesulfonate, monohydrate (9CI)
     INDEX NAME)
```

CM 1

CRN 79350-37-1

CMF C16 H15 N5 O7 S2

Absolute stereochemistry.

Double bond geometry as shown.

CM 2

CRN 75-75-2 CMF C H4 O3 S

RN 524925-13-1 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2Z)-(2-amino-4-thiazolyl)[(carboxymethoxy)imino]acetyl]amino]-3-ethenyl-8-oxo-, (6R,7R)-, monomethanesulfonate (9CI) (CA INDEX NAME)

CM 1

CRN 79350-37-1 CMF C16 H15 N5 O7 S2

CRN 75-75-2 C H4 O3 S CMF

IT 125110-14-7P, Cefixime trihydrate

RL: SPN (Synthetic preparation); PREP (Preparation) (process for the preparation of cefixime via alkylsulfonate or arylsulfonate salts)

125110-14-7 CAPLUS RN

5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2Z)-(2-amino-4-thiazolyl)[(carboxymethoxy)imino]acetyl]amino]-3ethenyl-8-oxo-, trihydrate, (6R,7R)- (9CI) (CA INDEX NAME)

## ●3 H<sub>2</sub>O

REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 16 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2001:880903 CAPLUS

DOCUMENT NUMBER: 137:125013

TITLE: Synthesis of cefdinir

AUTHOR(S): Lin, Gui-chun; Liu, Li; Ma, Ling-tai; Min, Ji-mei;

Zhang, Li-he

CORPORATE SOURCE: Natl. Res. Lab. Natural Biomimetic Drugs, Peking

Univ., Beijing, 100083, Peop. Rep. China

SOURCE: Hecheng Huaxue (2001), 9(5), 383-385

CODEN: HEHUE2; ISSN: 1005-1511

PUBLISHER: Hecheng Huaxue Bianjibu

DOCUMENT TYPE: Journal LANGUAGE: Chinese

OTHER SOURCE(S): CASREACT 137:125013

AB Cefdinir was synthesized via the condensation of 2-(2-aminothiazol-4-yl)-2-(Z)-(acetyinmino)acetyl chloride with 7-amino-3-vinyl-3-cephem-4-carboxylic acid. Under the optimization reaction conditions 60% total

yield was achieved.

IT 443874-49-5P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(synthesis of cefdinir)

RN 443874-49-5 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2Z)-[(acetyloxy)imino](2-amino-4-thiazolyl)acetyl]amino]-3-ethenyl-8-

oxo-, monohydrochloride, (6R,7R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

## ) HCl

ANSWER 17 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

2001:767504 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 135:303724

Preparation of 3-vinylcephem compound from protected TITLE:

compounds

Kameyama, Yutaka; Fukae, Kazuhiro INVENTOR(S): Ohtsuka Chemical Co., Ltd., Japan PATENT ASSIGNEE(S):

Jpn. Kokai Tokkyo Koho, 5 pp. SOURCE:

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT:

PATENT NO.	KIND DATE	APPLICATION NO.	DATE			
JP 2001294590	A2 2001102	3 JP 2000-111448	20000413			
WO 2001079211	A1 2001102	5 WO 2001-JP3182	20010413			
W: CN, KR						
RW: AT, BE, CH,	CY, DE, DK, ES	, FI, FR, GB, GR, IE,	IT, LU, MC, NL,			
PT, SE	•					
EP 1273587	A1 2003010	8 EP 2001-919924	20010413			
R: AT, BE, CH,		, GB, GR, IT, LI, LU, I	NL, SE, MC, PT,			
IE, FI, CY	•					
CN 1134445	B 2004011	.4 CN 2001-800920	20010413			
HK 1048112	A1 2004112	6 HK 2003-100146	20030107			
PRIORITY APPLN. INFO.:		JP 2000-111448	A 20000413			
		WO 2001-JP3182	W 20010413			
OTHER SOURCE(S):	CASREACT 135:3	03724; MARPAT 135:3037	24			

AB Cefdinir is prepared by treatment of protected 3-vinylcephem compds. I [R1-R3 = H, (un)substituted arylmethyl; R1 = R2 = R3 ≠ H] with perhalogenic acid and organic protonic acid in organic solvent. Thus, I (R1 = R3 = H, R2 = trityl) was treated with HClO4 and HCO2H at 30° for 1 h in CH2Cl2 to give 95% cefdinir.

IT 367267-68-3

RL: RCT (Reactant); RACT (Reactant or reagent)
 (preparation of 3-vinylcephem compound from protected compds.)

RN 367267-68-3 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,
7-[[(2Z)-(2-amino-4-thiazolyl)[(triphenylmethoxy)imino]acetyl]amino]-3ethenyl-8-oxo-, (6R,7R)-, compd. with 2-methyl-N-[(4methylphenyl)sulfonyl]propanamide (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 128454-32-0 CMF C33 H27 N5 O5 S2

Absolute stereochemistry.

Double bond geometry as shown.

CM 2

CRN 58821-27-5 CMF C11 H15 N O3 S

ANSWER 18 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN L9

ACCESSION NUMBER: 2001:748751 CAPLUS

DOCUMENT NUMBER: 136:25202

Spectrophotometric determination of cefixime TITLE:

trihydrate

Shankar, D. G.; Sushma, K.; Lakshmi, R. V.; Rao, Y. AUTHOR (S):

Srinivasa; Reddy, M. N.; Murthy, T. K.

Department of Pharmaceutical Sciences, Andhra CORPORATE SOURCE:

University, Visakhapatnam, 530 003, India

SOURCE: Asian Journal of Chemistry (2001), 13(4), 1649-1651

CODEN: AJCHEW; ISSN: 0970-7077

Asian Journal of Chemistry PUBLISHER: Journal

DOCUMENT TYPE: LANGUAGE: English

Two simple, sensitive and selective methods were developed for the AB determination

of cefixime in pure and pharmaceutical prepns. Method A is based on the formation of green colored chromogen by oxidative coupling reaction with 3-methyl-2-benzothiazolinone hydrazone (MBTH) and ferric chloride having absorption maximum at 620 nm, whereas method B is based on the reduction and complex formation with ferric chloride and 1,10-phenanthroline which exhibit maximum absorption at 510 nm. These methods obey Beer's law in the concentration range of 1 to 15 µg/mL and 0.2 to 6 µg/mL resp. The methods are statistically evaluated for accuracy and precision.

125110-14-7, Cefixime trihydrate ΙT

RL: ANT (Analyte); ANST (Analytical study)

(cefixime trihydrate determination in pure and pharmaceutical prepns. by spectrophotometry using Me benzothiazolinone hydrazone or ferric chloride and phenanthroline)

RN125110-14-7 CAPLUS

5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, CN

7-[[(2Z)-(2-amino-4-thiazolyl)[(carboxymethoxy)imino]acetyl]amino]-3-

ethenyl-8-oxo-, trihydrate, (6R,7R)- (9CI) (CA INDEX NAME)

## ●3 H<sub>2</sub>O

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 19 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

2001:713357 CAPLUS ACCESSION NUMBER:

135:272795 DOCUMENT NUMBER:

Process for preparing cephalosporin derivatives via a TITLE:

new thiazole compound

INVENTOR(S): Yoon, Dae Chul; Yoo, Seung Won; Shin, Dong Gyun; Lee,

Myoung Ki; Park, Mi Soon; Lee, Yoon Seok; Song, Yoon

Seok; Lee, Ju Cheol; Oh, Sang Mi

Hanmi Fine Chemicals Co. Ltd., S. Korea PATENT ASSIGNEE(S):

PCT Int. Appl., 16 pp. SOURCE:

CODEN: PIXXD2 DOCUMENT TYPE: Patent

English LANGUAGE:

FAMILY ACC. NUM. COUNT: 2

PATENT NO.					KIND DATE			APPLICATION NO.						DATE		
													-			
WO	2001	0707	49		A1	2001	.0927	WO	2001	-KR34	7		2	0010	307	
	W:	CN,	JP													
	RW:	ΑT,	BE,	CH,	CY,	DE, DK,	ES,	FI, F	R, GE	GR,	ΙE,	IT,	LU,	MC,	NL,	
		PT,	SE,	TR												
KR	2001	0921	30		Α	2001	1024	KR	2000	-1407	6		2	0000	320	
EP	1268	488			A1	2003	0102	EP	2001	-9125	31		2	0010	307	
	R:	AT,	BE,	CH,	DE,	DK, ES,	FR,	GB, G	R, IT	LI,	LU,	NL,	SE,	MC,	PT,	
		IE,	FI,	CY,	TR											
JР	2003	5281	05		T2	2003	0924	JP	2001	-5689	50		2	0010	307	
PRIORIT	Y APP	LN.	INFO	. :				KR	2000	-1407	6		A 2	0000	320	
								WO	2001	-KR34	7		W 2	0010	307	
OTHER SO	OURCE	(S):			CASI	REACT 13	5:27	2795; 1	MARPA	T 135	:272	795				

$$R^1$$
 $R^2$ 
 $O-C-CO_2H$ 
 $N$ 
 $C-CO-NH$ 
 $S$ 
 $CO_2H$ 
 $R^4$ 

AΒ A process for the preparation of cephalosporin antibiotics I (R1 and R2 = same or different and are H, alkyl group of 1-4 carbon atoms, cycloalkyl group of 3-5 carbon atoms; R4 = acetoxymethyl, pyridiniummethyl, vinyl; X = Cl, ... Br; acid in the acid addition salt = HCl, HBr, H2SO4, HClO4, formic, acetic, trifluoroacetic, propionic, methanesulfonic or benzenesulfonic acid) where... an acid addition salt of a crystalline aminothiazole compound II was acylated

via the reaction of a 7-aminocephalosporanic acid derivative III was accomplished. Thus cefixime trihydrate was produced in 87% yield via the reaction of 7-amino-3-vinyl-3-cephem-4-carboxylic acid in ClCH2Cl and N,O-bis(trimethylsilyl)acetamide followed by addition of (Z)-2-(2carboxymethoxyimino) -2-(2-thiazole-4-yl)acetylchloride HCl in Na hydrogen carbonate and iso-Pr ether and 6 N HCl. In this process using the aminothiazole II acylated by the 7-aminocephalosporanic acid derivative III in the indicated solvent, few or no byproducts were produced and the desired compound I could be directly obtained in high yield without the need for a deprotection step following acylation.

125110-14-7P, Cefixime trihydrate IT RL: IMF (Industrial manufacture); SPN (Synthetic preparation); PREP (Preparation)

(preparation of  $\beta$ -lactams via acylation with a new thiazole compound) 125110-14-7 CAPLUS

RN

5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2Z)-(2-amino-4-thiazolyl)[(carboxymethoxy)imino]acetyl]amino]-3ethenyl-8-oxo-, trihydrate, (6R,7R)- (9CI) (CA INDEX NAME)

## ●3 H<sub>2</sub>O

REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 20 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2001:688527 CAPLUS

DOCUMENT NUMBER: 136:58929

DOCUMENT NUMBER: 136:36929

TITLE: Reversed phase high performance liquid chromatographic

determination of cefixime in bulk drugs

AUTHOR(S): Gonzalez-Hernandez, Rolando; Nuevas-Paz, Lauro;

Soto-Mulet, Laritza; Lopez-Lopez, Miguel; Hoogmartens,

Joseph

CORPORATE SOURCE: Dpto. de Analisis, Centro de Quimica Farmaceutica,

Ciudad de La Habana, Cuba

SOURCE: Journal of Liquid Chromatography & Related

Technologies (2001), 24(15), 2315-2324

CODEN: JLCTFC; ISSN: 1082-6076

PUBLISHER: Marcel Dekker, Inc.

DOCUMENT TYPE: Journal LANGUAGE: English

AB A new technique for the quant. determination of cefixime trihydrate in bulk drugs

by HPLC was developed using simple reagents. Chosen conditions of anal.

were as follows: LiChrospher 100 RP - 18 (250 + 4 mm I.D.) column, mobile phase consisting of phosphate buffer pH 7.0 and MeCN (93:7, volume/volume), flow rate of 0,8 mL/min, loop of 20 μL and UV detection at 287 nm. The prospective validation of this technique showed that it is linear at 0.1-0.6 mg/mL (r = 0.9997), sensitive (0.3 %), precise (within-a-day repeatability, relative standard deviation = 1.0 %, day-to-day repeatability, relative standard deviation = 1.3 %), accurate and selective

repeatability relative standard deviation = 1.3 %), accurate and selective (cefixime can be determined in presence of its related compds.). The limits of detection and quantitation are 37 ng (0.3 %) and 128 ng (1.1 %), resp.,

relative to a 0.6 mg/mL solution IT 125110-14-7, Cefixime trihydrate

RL: ANT (Analyte); ANST (Analytical study)

(reversed phase high performance liquid chromatog. determination of cefixime in

bulk drugs)

RN 125110-14-7 CAPLUS

5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, CN7-[[(2Z)-(2-amino-4-thiazolyl)[(carboxymethoxy)imino]acetyl]amino]-3ethenyl-8-oxo-, trihydrate, (6R,7R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

#### H<sub>2</sub>O

THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 17

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

CAPLUS COPYRIGHT 2006 ACS on STN L9 ANSWER 21 OF 52

2001:564833 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER:

135:152367

TITLE:

Nitrate salts of antimicrobial agents

INVENTOR(S):

Del Soldato, Piero; Benedini, Francesca; Antognazza,

Patrizia

PATENT ASSIGNEE(S):

Nicox S.A., Fr.

SOURCE:

PCT Int. Appl., 105 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT NO.			KIND DATE		APPLICATION NO.						DATE						
						-											
WO	2001	0546	91		<b>A1</b>		2001	0802	1	WO 20	001-1	EP43	0		20010116		
	W:	AE.,	AL,	AU,	ŖΑ,	BB,	BG,	BR,	CA,	CN,	CR,	CU,	CZ,	DM,	EE,	GE,	HR,
		HU,	ID,	IL,	IN,	IS,	JP,	KΡ,	KR,	LC,	LK,	LR,	LT,	LV,	MA,	MG,	MK,
		MN,	MX,	NO,	NZ,	PL,	RO,	SG,	SI,	SK,	TR,	TT,	UA,	US,	UZ,	VN,	YU,
		ZA,	AM,	ΑZ,	BY,	KG,	ΚZ,	MD,	RU,	ТJ,	TM						
	RW:	GH,	GM,	ΚE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZW,	AT,	BE,	CH,	CY,
		DE,	DK,	ES,	FI,	FR,	GB,	GR,	ΙE,	IT,	LU,	MC,	NL,	PT,	SE,	TR,	BF,
		ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GW,	ML,	MR,	ΝE,	SN,	TD,	TG		
IT	1317	735	•		В1		2003	0715		IT 20	1-000	MI92			2	0000	126
CA	2397	754			AA		2001	0802	(	CA 2	001-	2397	754		2	0010	116
AU	2001	2001037308 A5 20010807		0807		AU 2	001-	3730	8		20010116						
BR	2001	0078	24	4 A 20021105			1105	BR 2001-7824						20010116			
EP	1253	924	A1 20021106			1106		EP 2	001-	9096	31		2	0010	116		

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,

IE, SI, LT, LV, FI, RO, MK, CY, AL, TR

JP 2003520814 T:2 20030708 JP 2001-554675 20010116 US 2003105066 A1 20030605 US 2002-181424 20020724

US 6794372 **B2** 20040921

PRIORITY APPLN. INFO.: IT 2000-MI92 Α 20000126

WO 2001-EP430 20010116 W

OTHER SOURCE(S): MARPAT 135:152367

Nitrate salts of antiviral, antifungal, and antibacterial agents such as acyclovir, tetracycline, etc. were prepared Growth inhibition of, e.g., an S. Aureus strain by title compds. was demonstrated.

352465-67-9P IT

> RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(nitrate salts of antimicrobial agents)

RN352465-67-9 CAPLUS

5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, CN7-[[(2Z)-(2-amino-4-thiazolyl)[(carboxymethoxy)imino]acetyl]amino]-3ethenyl-8-oxo-, (6R,7R)-, nitrate (9CI) (CA INDEX NAME)

CM 1

CRN 79350-37-1 CMF C16 H15 N5 O7 S2

Absolute stereochemistry. Double bond geometry as shown.

$$N = \frac{1}{2}$$
 $N = \frac{1}{2}$ 
 $N =$ 

CM 2

CRN 7697-37-2 CMF H N O3

REFERENCE COUNT: THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

Prepared by: Mary Hale @2-2507 Rem Bldg 1D86

L9 ANSWER 22 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2000:158096 CAPLUS

DOCUMENT NUMBER: 132:166060

TITLE: Preparation of crystalline salts of 7-[2-(2-aminothiazol-4-yl)-2-(tert-

butoxycarbonylmethoxyimino)acetamido]-3-vinyl-3-cephem-

4-carboxylic acid

INVENTOR(S): Decristoforo, Martin; Ludescher, Johannes; Miller,

Ludwig; Sturm, Hubert; Veit, Werner; Wolf, Siegfried

PATENT ASSIGNEE(S): Biochemie GmbH, Austria

SOURCE: Austrian, 10 pp. CODEN: AUXXAK

DOCUMENT TYPE: Patent

LANGUAGE: German FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
AT 405402	В	19990825	AT 1997-2058	19971204
AT 9702058	· A	19981215		
PRIORITY APPLN. INFO.:		•	AT 1997-2058	19971204
OTHER SOURCE(S):	MARPAT	132:166060	•	
GI ,				

AB Crystalline salts I · NR1R2R3 [R1 = R2 = R3 = Et; R1 = R2 = cyclohexyl, R3 = H; R1 = R2 = H, R3 = tert-octyl (CMe2CH2CMe3)] of 7-[2-(2-aminothiazol-4-yl)-2-(tert-butoxycarbonylmethoxyimino)acetamido]-3-vinyl-3-cephem-4-carboxylic acid (I) are prepared Thus, I·NCMe2CH2CMe3 was prepared via N-acylation of 7-amino-3-vinyl-3-cephem-4-carboxylic acid in EtOAc with 2-(aminothiazol-4-yl)-2-(tert-butoxycarbonylmethoxyimino)acetic acid S-mercaptobenzothiazolyl ester dimethylacetamide sulfate followed by mixing with Me3CCH2CMe2NH2 in AcOEt.

IT 210702-13-9P 210702-14-0P 210702-15-1P 258871-56-6P 258871-57-7P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of crystalline salts of a 3-vinyl-3-cephem-4-carboxylic acid derivative)

RN 210702-13-9 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,
7-[[(2Z)-(2-amino-4-thiazolyl)[[2-(1,1-dimethylethoxy)-2-oxoethoxy]imino]acetyl]amino]-3-ethenyl-8-oxo-, (6R,7R)-, compd. with 2,4,4-trimethyl-2-pentanamine (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 79368-92-6

Prepared by: Mary Hale @2-2507 Rem Bldg 1D86

CMF C20 H23 N5 O7 S2

Absolute stereochemistry.

Double bond geometry as shown.

CM 2

CRN 107-45-9 CMF C8 H19 N

$$\begin{array}{c} \text{NH}_2 \\ | \\ \text{Me-C-CH}_2\text{--CMe}_3 \\ | \\ \text{Me} \end{array}$$

RN 210702-14-0 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,

7-[[(2Z)-(2-amino-4-thiazolyl)[[2-(1,1-dimethylethoxy)-2-oxoethoxy]imino]acetyl]amino]-3-ethenyl-8-oxo-, (6R,7R)-, compd. with

N, N-diethylethanamine (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 79368-92-6 CMF C20 H23 N5 O7 S2

CRN 121-44-8 CMF C6 H15 N

RN 210702-15-1 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2Z)-(2-amino-4-thiazolyl)[[2-(1,1-dimethylethoxy)-2-oxoethoxy]imino]acetyl]amino]-3-ethenyl-8-oxo-, (6R,7R)-, compd. with N-cyclohexylcyclohexanamine (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 79368-92-6 CMF C20 H23 N5 O7 S2

Page 50

CM 2

CRN 101-83-7 CMF C12 H23 N

RN 258871-56-6 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,
7-[[(2E)-2-(2-amino-4-thiazolyl)-2-[[2-(1,1-dimethylethoxy)-2-oxoethoxy]imino]ethyl]amino]-3-ethenyl-8-oxo-, (6R,7R)-, sulfate (1:1),
monohydrate (9CI) (CA INDEX NAME)

CM 1

CRN 79368-92-6 CMF C20 H23 N5 O7 S2

Absolute stereochemistry.

Double bond geometry as shown.

CM 2

CRN 7664-93-9 CMF H2 O4 S

RN 258871-57-7 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,
7-[[(2E)(2-amino-4-thiazolyl)[[2-(1,1-dimethylethoxy)-2-oxoethoxy]imino]acetyl]amino]-3-ethenyl-8-oxo-, trihydrate, (6R,7R)- (9CI)

## (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

# ●3 H<sub>2</sub>O

L9 ANSWER 23 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1999:672823 CAPLUS

DOCUMENT NUMBER:

131:286331

TITLE:

Process for producing a cephem compound by

deprotection using phenols or phenols and protonic

acids

INVENTOR(S):

Kameyama, Yutaka

PATENT ASSIGNEE(S):

Otsuka Kagaku Kabushiki Kaisha, Japan

SOURCE: PCT Int. Appl., 15 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT NO.	KIND DATE	APPLICATION NO.	DATE			
WO 9952913	A1 19991021	WO 1999-JP1942	19990413			
W: AE, AL, AM,	AT, AU, AZ, BA,	BB, BG, BR, BY, CA,	CH, CN, CU, CZ,			
DE, DK, EE,	ES, FI, GB, GD,	GE, GH, GM, HR, HU,	ID, IL, IN, IS,			
JP, KE, KG,	KP, KR, KZ, LC,	LK, LR, LS, LT, LU,	LV, MD, MG, MK,			
MN, MW, MX,	NO, NZ, PL, PT,	RO, RU, SD, SE, SG,	SI, SK, SL, TJ,			
TM, TR, TT,	UA, UG, US, UZ,	VN, YU, ZA, ZW, AM,	AZ, BY, KG, KZ,			
MD, RU, TJ,	TM					
RW: GH, GM, KE,	LS, MW, SD, SL,	SZ, UG, ZW, AT, BE,	CH, CY, DE, DK,			
ES, FI, FR,	GB, GR, IE, IT,	LU, MC, NL, PT, SE,	BF, BJ, CF, CG,			
CI, CM, GA,	GN, GW, ML, MR,	NE, SN, TD, TG				
· AU 9934432	A1 19991101	AU 1999-34432	19990413			
PRIORITY APPLN. INFO.:		JP 1998-121888	A 19980414			
		WO 1999-JP1942	W 19990413			
OTHER SOURCE(S):	CASREACT 131:28	6331; MARPAT 131:2863	31			

$$C - CO - NH$$
 $C - CO - NH$ 
 $C - CO - NH$ 

AB Cephem compound I is prepared by deprotecting II [R1 = H, CHO, trityl group containing electron donating groups on the Ph rings; R2 = tert-Bu, naphthylmethyl, anthrylmethyl, benzyl group containing electron donating groups on the Ph ring, benzhydryl group containing electron donating groups on the Ph rings; R3 = naphthylmethyl, anthrylmethyl, benzyl group containing electron donating groups on the Ph ring] with phenols alone, or with a combination of phenols and protonic acids. Thus, II [R1 = H, R2 = t-Bu, R3 = CH2-C6H4-OMe-p] was stirred with p-toluenesulfonic acid and m-cresol at room temperature for 3 h to give 98.6% I.

IT 202843-53-6P

RL: IMF (Industrial manufacture); SPN (Synthetic preparation); PREP (Preparation)

(process for producing cephem compound by deprotection using phenols or phenols and protonic acids)

RN 202843-53-6 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2Z)-(2-amino-4-thiazolyl)[(carboxymethoxy)imino]acetyl]amino]-3-ethenyl-8-oxo-, (6R,7R)-, mono(4-methylbenzenesulfonate) (9CI) (CA INDEX NAME)

CM 1

CRN 79350-37-1 CMF C16 H15 N5 O7 S2

Absolute stereochemistry.

Double bond geometry as shown.

CM 2

CRN 104-15-4 C7 H8 O3 S CMF

REFERENCE COUNT:

THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS 10 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 24 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1999:659390 CAPLUS

DOCUMENT NUMBER:

131:286328

TITLE:

Process for purification of cefixime, a cephalosporin

derivative

INVENTOR(S):

Decristoforo, Martin; Ludescher, Johannes; Sturm,

Hubert

PATENT ASSIGNEE(S):

Biochemie G.m.b.H., Austria

SOURCE:

PCT Int. Appl., 18 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

						KIND DATE			APPLICATION NO.						DATE				
			;																
											WO 1	.999-1	EP22:	22			19990	331	
	WO	9951																	
		W:															, CU,		
																	, IN,		
		:	JP,	KE,	KG,	ΚP,	KR,	ΚZ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	MD	, MG,	MK,	-
			MN,	MW,	MX,	NO,	NZ,	PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK	, SL,	TJ;	
			TM,	TR,	TT,	UA,	UG,	US,	UΖ,	VN,	YU,	ZA,	·ZW,	ΑM,	ΑZ,	BY	, KG,	ΚZ,	
			MD,	RU,	ТJ,	TM							-						
		RW:	GH,	GM,	KE,	LS,	MW,	SD,	SL,	SZ,	ŪG,	ZW,	ΑT,	BE.,	CH,	CY	, DE,	DK,	
			ES,	FI,	FR,	GB,	GR,	ΙE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	BJ	, CF,	CG,	
			CI,	CM,	GΑ,	GN,	GW,	ML,	MR,	NE,	SN,	TD,	TG						
	AT	9800	575			Α		2000	0115		AT 1	998-	575				19980	402	
	AΤ	4067	73			В		2000	0825						•				
	CA	23264	441			AA		1999	1014	4	CA 1	999-	2326	441		:	19990	331	
																	19990		
		99098										999-					19990		
	EΡ	10682	211			A2		2001	0117		EP 1	999-	9179	38	•	:	19990	331	
																	, PT,		
			SI,				•	•	•		•	•	·	·	·			•	
	TR	2000						2001	0221	٠ .	TR 2	2000-	2000	02838	3		19990	331	
	.TD	20021	5106	9.4		ΤЭ		2002	0409	1	JP 2	2000-	5423	28			19990	331	
	CN	11344	146		•	. в		2004	0114		CN 1	999-	3046	20			19990	331	
	ZA	20000	0048	99		A		2001	1014		ZA 2	2000-4	1899				20000	914	
	US	20032	2080	55		A1	•					2002-					20020		
		6825																	
PRTO		APP									<b>ΔΤ 1</b>	998-	575			Α.	19980	402	
11.10.										,	WO 1	1999-1	EP22	22	1	W .	19990		
																	20000		

$$H_2N$$
 $N$ 
 $CCO HN$ 
 $N$ 
 $CH = CH_2$ 
 $OCH_2CO_2R$ 
 $CO_2H$ 
 $I$ 

$$H_2N$$
 $H_2N$ 
 $H_2N$ 

AB A process for the production and purification of a cephalosporin derivative I (R =

alkyl or aryl where the amine group attached to the thiazolyl ring is free or protected) comprising reacting II in free form, protected form or a salt with a compound III (R = defined above and the amine group attached to the thiazolyl ring is free or protected) was accomplished. Thus cefixime I (R = Me) as the H2NCMe2CH2CMe3 salt was prepared via the reaction of 2-(2-amino-4-thiazolyl)-(Z)-2-(methoxycarbonylmethoxyimino)acetic acid and 2,2'-benzothiazolyl disulfide and the product obtained was further reacted with 7-amino-3-vinylceph-3-em-4-carboxylic acid followed by tert-octylamine.

IT 246035-37-0P

RL: IMF (Industrial manufacture); PUR (Purification or recovery); RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and purification the  $\beta$ -lactam cefixime)

RN 246035-37-0 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,
7-[[(2Z)-(2-amino-4-thiazolyl)[(2-methoxy-2-oxoethoxy)imino]acetyl]amino]3-ethenyl-8-oxo-, (6R,7R)-, compd. with 2,4,4-trimethyl-2-pentanamine
(1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 88621-01-6 CMF C17 H17 N5 O7 S2

MeO 
$$\sqrt{Z}$$
  $\sqrt{Z}$   $\sqrt{Z$ 

CRN 107-45-9 CMF C8 H19 N

$$\begin{array}{c} \text{NH}_2 \\ | \\ \text{Me-C-CH}_2\text{--CMe}_3 \\ | \\ \text{Me} \end{array}$$

IT 125110-14-7P

RL: IMF (Industrial manufacture); PUR (Purification or recovery); SPN (Synthetic preparation); PREP (Preparation) (preparation and purification the  $\beta$ -lactam cefixime)

RN 125110-14-7 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,
7-[[(2Z)-(2-amino-4-thiazolyl)[(carboxymethoxy)imino]acetyl]amino]-3-ethenyl-8-oxo-, trihydrate, (6R,7R)- (9CI) (CA INDEX NAME)

## ●3 H<sub>2</sub>O

L9 ANSWER 25 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1998:710723 CAPLUS

DOCUMENT NUMBER: 130:66294

TITLE: An effective and convenient esterification of

cephalosporin derivatives by using quaternary ammonium

salts as catalysts

AUTHOR(S): Lee, Hong Woo; Kang, Tae Won; Kim, Eung-Nam; Shin,

Jaewook; Cha, Kyung Hoi; Cho, Dong Ock; Choi, Nam Hee;

Kim, Jung-Woo; Hong, Chung, II

CORPORATE SOURCE: Research Institute, Chong Kun Dang Corp., Seoul,

152-600, S. Korea

SOURCE: Synthetic Communications (1998), 28(23), 4345-4354

CODEN: SYNCAV; ISSN: 0039-7911

PUBLISHER:

Marcel Dekker, Inc.

DOCUMENT TYPE:

Journal English

LANGUAGE:

OTHER SOURCE(S):

CASREACT 130:66294

AB A method for preparing cephalosporin derivs. by reacting cephalosporin alkaline metal salts with organic halide in the presence of quaternary ammonium salts catalyst is disclosed.  $\Delta 3$  To  $\Delta 2$  isomerization, a side reaction commonly reported in preparation of cephalosporin derivs., was successfully eliminated. The desired  $\Delta 3$  was obtained as a sole

IT 79350-44-0

RL: RCT (Reactant); RACT (Reactant or reagent)

(esterification of cephalosporin derivs. via quaternary ammonium salt catalysis)

RN 79350-44-0 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,

7-[[(2Z)-(2-amino-4-thiazolyl)(methoxyimino)acetyl]amino]-3-ethenyl-8-oxo-, monosodium salt, (6R,7R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

product in the reaction.

Na

REFERENCE COUNT:

13 THERE ARE 13 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 26 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1998:509198 CAPLUS

DOCUMENT NUMBER:

129:136023

TITLE:

Preparation of cefixime from aminovinylcephemcarboxylate and

(aminothiazolyl) (carboxymethoxyimino) acetic acid

derivatives

INVENTOR(S):

Ludescher, Johannes; Miller, Ludwig; Sturm, Hubert; Veit, Werner; Decristoforo, Martin; Wolf, Siegfried

PATENT ASSIGNEE(S):

Biochemie G.m.b.H., Austria PCT Int. Appl., 31 pp.

SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PAT	PATENT NO.							DATE APPLICATION NO.							DATE			
						-	<u></u>								-			
WO	9831	685			A1		1998	0723		WO 1	998-	EP19	0		1:	9980	114	
	W:	AL,	AM,	ΑT,	AU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CU,	CZ,	DE,	
		DK,	EE,	ES,	FI,	GB,	GE,	GH,	GM,	GW,	HU,	ID,	IL,	IS,	JP,	ΚE,	KG,	
		KP,	KR,	ΚZ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	MD,	MG,	MK,	MN,	MW,	MX,	
		NO,	NZ,	PĹ,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TR,	TT,	
		UA,	UG;	US,	UΖ,	VN,	YU,	ZW,	AM,	AZ,	BY,	KG,	KZ,	MD,	RU,	ТJ,	TM	
	RW:	GH,	GM,	KE,	LS,	MW,	SD,	SZ,	UG,	ZW,	AT,	BE,	CH,	DE,	DK,	ES,	FI,	
		FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,	CF,	CG,	CI,	CM,	
		GA,	GN,	ML,	MR,	NE,	SN,	TD,	TG									
ΑT	9700	061			Α		1998	0615		AT 1	997-	61			1	9970	116	
AT	4047	26		7	В		1999	0225										
ΑT	9700	062			Α		1998	0615		AT 1	997-	62			1	9970	116	
AΤ	4047	27			В		1999	0225										
TW	5380	45			В		2003	0621		TW 1	998-	8710	0131		1	9980	107	
AU	9866	141			A1		1998	0807		AU 1	998-	6614	1		1	9980	114	
ΕP	9682	14			A1		2000	0105		EP 1	998-	9079	45		1	9980	114	
ΕP	9682	14	•		В1		2004	0407										

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R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, FI
     JP 2000514090
                         T2
                               20001024
                                            JP 1998-533299
                                                                  19980114
     AT 263774
                         E
                               20040415
                                           AT 1998-907945
                                                                  19980114
     ES 2219874
                         Т3
                               20041201
                                           ES 1998-907945
                                                                  19980114
                         B1
                                           US 1999-341542
     US 6313289
                               20011106
                                                                  19990804
                         A1
                               20050128
                                           HK 2000-104089
     HK 1024698
                                                                  20000704
     JP 2004155793
                         A2
                               20040603
                                           JP 2004-10146
                                                                  20040119
PRIORITY APPLN. INFO.:
                                           AT 1997-61
                                                               A 19970116
                                           AT 1997-62
                                                               A 19970116
                                            EP 1998-907945
                                                               A 19980114
                                            JP 1998-533299
                                                               A3 19980114
                                            WO 1998-EP190
                                                               W 19980114
OTHER SOURCE(S):
                       CASREACT 129:136023; MARPAT 129:136023
GT
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\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

AB The process of preparing cefixime [I; R = NH2, R1 = R2 = H] involves reaction of II [R7 = H, alkyl, cycloalkyl, alkylaryl, aryl, aralkyl, silyl; R5, R6 = H, leaving group] with a 2-(aminothiazol-4-yl)-2- (carboxymethoxyimino)acetic acid derivative [III; R9 = alkyl, cycloalkyl, alkylaryl, aryl, aralkyl; R10 = H; R11 = H, silyl, acyl], reacting the resulting I [R = NR10R11, R1 = R7, R2 = R9] (IV) with NR1R2R3 [R1, R2, R3 = H, alkyl, cycloalkyl, alkylaryl, aryl, aralkyl], treating the resulting crystalline IV.NR1R2R3 with H2SO4, and decomposing the resulting cefixime sulfate.

Thus, III [R9 = t-Bu, R10 = R11 = H].MeCONMe2 (preparation given) was reacted with II [R5 = R6 = R7 = H] in aqueous EtOAc containing Et3N and the product treated with H3PO4 and then tert-octylamine to give I [R = NH2, R1 = H, R2 = tBu].tert-octylamine, which was treated with H2SO4 in MeCN containing HCOOH to give cefixime addition salt with sulfuric acid, which in water was treated with NH3 to give cefixime of 99% purity.

IT 210702-13-9P 210702-14-0P 210702-15-1P 210702-16-2P

RL: IMF (Industrial manufacture); RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (preparation of cefixime from aminovinylcephemcarboxylate and (aminothiazolyl) (carboxymethoxyimino) acetic acid derivs.)

RN 210702-13-9 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,
7-[[(2Z)-(2-amino-4-thiazolyl)[[2-(1,1-dimethylethoxy)-2-oxoethoxy]imino]acetyl]amino]-3-ethenyl-8-oxo-, (6R,7R)-, compd. with 2,4,4-trimethyl-2-pentanamine (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 79368-92-6 CMF C20 H23 N5 O7 S2

CRN 107-45-9 CMF C8 H19 N

$$\begin{array}{c} \text{NH}_2 \\ | \\ \text{Me-C-CH}_2\text{--CMe}_3 \\ | \\ \text{Me} \end{array}$$

RN 210702-14-0 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,

7-[[(2Z)-(2-amino-4-thiazolyl)[[2-(1,1-dimethylethoxy)-2-oxoethoxy]imino]acetyl]amino]-3-ethenyl-8-oxo-, (6R,7R)-, compd. with N,N-diethylethanamine (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 79368-92-6 CMF C20 H23 N5 O7 S2

CRN 121-44-8 CMF C6 H15 N

RN 210702-15-1 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2Z)-(2-amino-4-thiazolyl)[[2-(1,1-dimethylethoxy)-2-oxoethoxy]imino]acetyl]amino]-3-ethenyl-8-oxo-, (6R,7R)-, compd. with N-cyclohexylcyclohexanamine (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 79368-92-6 CMF C20 H23 N5 O7 S2

Absolute stereochemistry. Double bond geometry as shown.

CM 2

CRN 101-83-7 CMF C12 H23 N

RN 210702-16-2 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2Z)-(2-amino-4-thiazolyl)[(carboxymethoxy)imino]acetyl]amino]-3-ethenyl-8-oxo-, (6R,7R)-, sulfate (1:1) (9CI) (CA INDEX NAME)

Prepared by: Mary Hale @2-2507 Rem Bldg 1D86

CRN 79350-37-1

CMF C16 H15 N5 O7 S2

Absolute stereochemistry.

Double bond geometry as shown.

CM 2

CRN 7664-93-9 CMF H2 O4 S

REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 27 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1998:126256 CAPLUS

DOCUMENT NUMBER:

128:167306 .

TITLE:

Purification of cefixime via amine salts

INVENTOR(S):

Miller, Ludwig; Sturm, Hubert

PATENT ASSIGNEE(S):

Biochemie G.m.b.H., Austria; Miller, Ludwig; Sturm,

Hubert

SOURCE:

PCT Int. Appl., 16 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 3

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9806723	A1	19980219	WO 1997-EP4439	19970813
W: AL, AM, AT,	AU, AZ	, BA, BB, BG	, BR, BY, CA, CH, CN,	CU, CZ, DE,

DK, EE, ES, FI, GB, GE, HU, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, TJ, TM, TR, TT, UA, UG, US, UZ, VN, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG AT 9601468 Α 19980215 AT 1996-1468 19960814 AT 404251 В 19981027 AU 9746168 **A1** 19980306 AU 1997-46168 19970813 PRIORITY APPLN. INFO.: AT 1996-1468 Δ 19960814 WO 1997-EP4439 W 19970813

AB Cefixime in form of a salt with dicyclohexylamine, e.g. a bis-dicyclohexylammonium salt, was prepared in a process for purification of cefixime. Thus, impure cefixime-trihydrate was treated with dicyclohexylamine in acetone and water to give cefixime-bis-dicyclohexylamine salt, which was mixed with water and activated carbon and the pH adjusted to 2.5 by addition of sulfuric acid to give cefixime trihydrate.

IT 125110-14-7P 202843-54-7P

RL: PUR (Purification or recovery); RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (purification of cefixime via amine salts)

RN 125110-14-7 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2Z)-(2-amino-4-thiazolyl)[(carboxymethoxy)imino]acetyl]amino]-3-ethenyl-8-oxo-, trihydrate, (6R,7R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

#### ●3 H<sub>2</sub>O

RN 202843-54-7 CAPLUS
CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,
7-[[(2-amino-4-thiazolyl)[(carboxymethoxy)imino]acetyl]amino]-3-ethenyl-8oxo-, [6R-[6α,7β(Z)]]-, compd. with N-cyclohexylcyclohexanamine
(1:2) (9CI) (CA INDEX NAME)

CM 1

Page 63

CRN 79350-37-1 CMF C16 H15 N5 O7 S2

Absolute stereochemistry.

Double bond geometry as shown.

CM 2

CRN 101-83-7 CMF C12 H23 N

IT 202843-53-6P

RL: SPN (Synthetic preparation); PREP (Preparation)
 (purification of cefixime via amine salts)

RN 202843-53-6 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,
7-[[(2Z)-(2-amino-4-thiazolyl)[(carboxymethoxy)imino]acetyl]amino]-3ethenyl-8-oxo-, (6R,7R)-, mono(4-methylbenzenesulfonate) (9CI) (CA INDEX NAME)

CM 1

CRN 79350-37-1 CMF C16 H15 N5 O7 S2

CRN 104-15-4 CMF C7 H8 O3 S

REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 28 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1997:547291 CAPLUS

DOCUMENT NUMBER: 127:149040

TITLE: Process for preparation of cefdinir

INVENTOR(S): Lee, Gwan Sun; Chang, Young Kil; Chun, Jong Pil; Koh,

Joon Hyung

PATENT ASSIGNEE(S): Hanmi Pharmaceutical Co., Ltd., S. Korea; Lee, Gwan

Sun; Chang, Young Kil; Chun, Jong Pil; Koh, Joon Hyung

SOURCE: PCT Int. Appl., 26 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT NO.	KIND DATE	APPLICATION NO.	DATE
WO 9724358	A1 19970710	WO 1996-KR250	19961226
W: JP, US			
RW: AT, BE, CH,	DE, DK, ES, FI,	FR, GB, GR, IE, IT, LU,	, MC, NL, PT, SE
KR 174432	B1 19990218	KR 1995-58694	19951227
KR 174431	B1 19990218	KR 1995-58695	19951227
EP 874853	A1 19981104	EP 1996-943357	19961226
EP 874853	B1 20020605		
R: AT, BE, CH,	DE, DK, ES, FR,	GB, GR, IT, LI, LU, NL,	, SE, MC, PT,
IE, FI			

JP 2000502700	Т2	20000307	JP	1997-524230		19961226
AT 218572	Ε.	20020615	AT	1996-943357		19961226
PT 874853	T	20020930	PT	1996-943357		19961226
ES 2175167	Т3	20021116	ES	1996-943357		19961226
US 6093814	A	20000725	ÜS	1998-68719		19980518
PRIORITY APPLN. INFO.:			KR	1995-58694	Α	19951227
			KR	1995-58695	Α	19951227
			WO	1996-KR250	W	19961226
OTHER SOURCE(S):	CASRE	ACT 127:14904	10; 1	MARPAT 127:149040	)	
GI .						•

Cefdinir I (R = H), a cephalosporin antibiotic, was prepared in an excellent color and purity and with a good yield. Cefdinir was prepared by N-acylation of 7-amino-3-vinyl-3-cephem-4-carboxylic acid with thio ester II (Z = 2-benzothiazolylthio) and crystallization of the resulting ester with 4-MeC6H4SO3H and Me2NCOMe to form crystals of I (R = CPh3).4-MeC6H4SO3H.2Me2NCOMe, which were then converted to cefdinir with the use of formic acid. Formation of the cefdinir amide linkage was also accomplished starting from phosphoryl ester II [Z = OP(O)(OEt)2].

IT 193402-46-9P

RI: IMF (Industrial manufacture): PRP (Properties): RCT (Reactant): SPN

RN 193402-46-9 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2-amino-4-thiazolyl)[(triphenylmethoxy)imino]acetyl]amino]-3-ethenyl-8-oxo-, [6R-[6 $\alpha$ ,7 $\beta$ (Z)]]-, mono(4-methylbenzenesulfonate), compd. with N,N-dimethylacetamide (1:2) (9CI) (CA INDEX NAME)

CM 1 CRN 127-

CRN 127-19-5 CMF C4 H9 N O

CRN 193402-45-8

C33 H27 N5 O5 S2 . C7 H8 O3 S CMF

> CM 3

CRN 128454-32-0

CMF C33 H27 N5 O5 S2

Absolute stereochemistry. Double bond geometry as shown.

CM

CRN 104-15-4 CMF C7 H8 O3 S

L9 ANSWER 29 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1997:253486 CAPLUS

DOCUMENT NUMBER: TITLE:

Studies on new catechol containing cephalosporins. III. Synthesis and structure-activity relationships of

cephalosporins having a pyridone moiety at the C-7

position

126:277312

AUTHOR(S):

Choi, Kyung Il; Cha, Joo Hwan; Pae, Ae Nim; Cho, Yong Seo; Koh, Hun Yeong; Chang, Moon Ho; Kang, Han-Young;

Prepared by: Mary Hale @2-2507 Rem Bldg 1D86

Page 67

Chung, Bong Young

CORPORATE SOURCE: Division of Applied Science, Korea Institute of

Science and Technology, Seoul, 130-650, S. Korea

SOURCE: Journal of Antibiotics (1997), 50(3), 279-282

CODEN: JANTAJ; ISSN: 0021-8820

PUBLISHER: Japan Antibiotics Research Association

DOCUMENT TYPE: Journal LANGUAGE: English

GΙ

AB Cephalosporins I [Q = H, Cl, CH:CH2,CH2OAc, 1-methyl-5-tetrazolylthiomethyl, R = CO2Na; Q = 1-R1-pyridinium-4-ylthiomethyl, R1 = Et, CH2CH2OH, CH2CO2-, NHMe, R = CO2-] were prepared I all exhibit good antibacterial activity against both gram-pos. and gram-neg. bacteria, especially

Ι

Pseudomonas aeruginosa.

IT 189017-35-4P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation of bactericidal pyridylisoxazolylmethoxyiminoacetamidocephalosporins)

RN 189017-35-4 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2-amino-4-thiazolyl)[[[3-(1,4-dihydro-1,5-dihydroxy-4-oxo-2-pyridinyl)-5-isoxazolyl]methoxy]imino]acetyl]amino]-3-ethenyl-8-oxo-, disodium salt,  $[6R-[6\alpha,7\beta(Z)]]$ - (9CI) (CA INDEX NAME)

#### 2 Na

REFERENCE COUNT: 7 THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 30 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

1996:110431 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER:

124:145747

Process for the preparation of trihydrated cefixime TITLE:

Picornell Dardes, Carlos INVENTOR(S):

PATENT ASSIGNEE(S): Marcham Trading Investment Ltd., Ire.

SOURCE: PCT Int. Appl., 24 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent French LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.									APPL	ICAT	ION I	NO.	DATE				
						_									-		
WC	9533	753			A1		1995	1214	1	WO 1	995-	EP17	59		1	9950	510
	W:	AM,	AT,	AU,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CZ,	DE,	DK,	ES,	FI,	GB,
		GE,	HU,	JP,	KE,	KG,	ΚP,	KR,	ΚZ,	LK,	LT,	LU,	LV,	MD,	MG,	MN,	MW,
		NO,	NZ,	PL,	PT,	RO,	RU,	SD,	SE,	SI,	SK,	TJ,	TT,	ŲΑ,	US,	UZ,	VN
	RW:	ΚE,	MW,	SD,	SZ,	ŪĠ,	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙE,	IT,
		LU,	MC,	NL,	PT,	SE,	BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	ML,	MR,	NE,
		SN,	TD,	TG													
CH	6883	19			Α		1997	0731	(	CH 1	994-	1751			1	9940	603
AU	9526	702			<b>A1</b>		1996	0104	i	AU 1	995-	2670	2		1	9950	510
EF	7630	43			<b>A1</b>		1997	0319	1	EP 1	995-	9217	39		1	9950	510
EF	7630	43			В1		1998	0923									
	R:	AT,	BE,	CH,	DE,	DK,	FR,	GB,	GR,	ΙE,	IT,	LI,	LU,	NL,	PT,	SE	
ΓA	1714	54			E		1998	1015	i	AT 1	995-	9217	39		1	9950	510
ES	2120	911			<b>A1</b>		1998	1101	]	ES 1	996-	5000	4		1	9960	202
ES	2120	911			B1		1999	0701									
PRIORIT	Y APP	LN.	INFO	. :					(	CH 1	994-	1751			A 1	9940	603
											995-						
OTHER SOURCE(S):					CASREACT 124:14!				5747								

GI

The invention relates to a process for the preparation of trihydrated cefixime (I) by reacting a functional derivative of N-protected (Z)-2-(2-aminothiazol-4-yl)-2-(tert-butoxycarbonylmethoxyimino)acetic acid with tert-Bu 7-amino-3-vinyl-3-cephem-4-carboxylate, or one of the salts thereof and, after removal of the protection group from the product thus obtained, by treating the product of the reaction with aluminum trichloride and anisole. This new process is carried out by using the new intermediate  $7\beta$ -[(Z)-2-(2-aminothiazol-4-yl)-2-(tert-butoxycarbonylmethoxyimino)ac etamido]-3-vinyl-3-cephem-4-carboxylate of tert-Bu, optionally N-protected on the thiazolic amine.

Ι

IT 125110-14-7P, Cefixime trihydrate

RL: SPN (Synthetic preparation); PREP (Preparation) (process for the preparation of trihydrated cefixime)

RN 125110-14-7 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2Z)-(2-amino-4-thiazolyl)[(carboxymethoxy)imino]acetyl]amino]-3-ethenyl-8-oxo-, trihydrate, (6R,7R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

●3 H<sub>2</sub>O

L9 ANSWER 31 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: DOCUMENT NUMBER:

1995:968353 CAPLUS

TITLE:

124:116911
Studies on new catechol containing cephalosporins. II.
Synthesis and structure-activity relationships of cephalosporins having a catechol moiety at the C-7 position

Page 70

AUTHOR(S): Choi, Kyung Il; Cha, Joo Hwan; Pae, Ae Nim; Cho, Y ong

Seo; Kang, Han-Young; Koh, Hun Yeong; Chang, Moon Ho

CORPORATE SOURCE: Div. Applied Science, Korea Inst. Science Technology,

Seoul, 130-650, S. Korea

SOURCE: Journal of Antibiotics (1995), 48(11), 1375-7

CODEN: JANTAJ; ISSN: 0021-8820

PUBLISHER: Japan Antibiotics Research Association

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 124:116911

GI

$$R^2 = \sqrt{N^+ \text{ Et}}$$
  $R^3 = \sqrt{N^+ \text{ CH}_2\text{CO}_2}$ 

AB We wish to report the synthesis and structure-activity relationship of cephalosporins, e.g. I (X = H, Cl; R = H, CH2OAc, CH:CH2, CH2SR1, CH2SR2, CH2SR3), having a catechol moiety at the C-7 position.

IT 172699-04-6P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(synthesis and structure-activity relationships of catechol contg cephalosporins)

RN 172699-04-6 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,
7-[[(2-amino-4-thiazolyl)[[[3-(2,5-dichloro-3,4-dihydroxyphenyl)-5-isoxazolyl]methoxy]imino]acetyl]amino]-3-ethenyl-8-oxo-, monosodium salt,
(6R-trans)- (9CI) (CA INDEX NAME)

Na

L9 ANSWER 32 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1994:700665 CAPLUS

DOCUMENT NUMBER: 121:300665

TITLE: Preparation of cephem derivatives as bactericides

INVENTOR(S): Moon, Ho Chang; Kang, Han Young; Ko, Hoon Young

PATENT ASSIGNEE(S): Korea Institute of Science and Technology, Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 18 pp.

Patent

CODEN: JKXXAF

DOCUMENT TYPE:

LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
			TD 1003 185053		,
JP 06179683	A2	19940628	JP 1993-175253		19930715
JP 2549494	B2	19961030	KD 1002 12641		10000715
KR 9508318	B1	19950727	KR 1992-12641	7\	19920715 19920715
PRIORITY APPLN. INFO.:	MADDAM	101 200665	KR 1992-12641	A	19920/15
OTHER SOURCE(S):	MARPAT	121:300665			
GI ·					

# \* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

AB The title compds. I [R1 = H, protecting group; R2 = H, salt-forming atom, etc.; R3, R4 = H, protecting group; Q = H, halo, etc.] are prepared Cephem (Z)-II (preparation given) in vitro showed MICs of 0.098, <0.002, and 0.098 μg/mL against Streptococcus pyogenes A308, Escherichia coli DC 2, and Pseudomonas aeruginosa 9027, resp. The antibacterial activities of 7 compds. of this invention are given in this document.

IT 159048-25-6P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of cephem bactericides)

Page 72

RN 159048-25-6 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,

7-[[(2-amino-4-thiazolyl)[[[3-(1,4-dihydro-1,5-dihydroxy-4-oxo-2-pyridinyl)-5-isoxazolyl]methoxy]imino]acetyl]amino]-3-ethenyl-8-oxo-,

monosodium salt,  $[6R-[6\alpha,7\beta(Z)]]-(9CI)$  (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

● Na

L9 ANSWER 33 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1994:409027 CAPLUS

DOCUMENT NUMBER: 121:9027

TITLE: Preparation of (pyridiniomethyl)cephemcarboxylates and

analogs as antibacterial agents

INVENTOR(S): Takamura, Norio; Saito, Kunio; Matsushita, Tadahiro;

Yamaguchi, Totaro

PATENT ASSIGNEE(S): Tanabe Seiyaku Co, Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 47 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

LANGUAGE: J FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE	
JP 05202062	A2	19930810	JP 1992-53045	19920127	
PRIORITY APPLN. INFO.:			JP 1992-53045	19920127	
OTHER SOURCE(S):	MARPAT	121:9027			
GI					

The title compds. I [R1 = (protected) amino; R2 = (protected) OH, alkoxy; R3 = (protected) carboxyl; R4 = H, alkyl, CH2R41, etc.; R41 = nucleophilic moiety; R5 = (protected) carboxyl, CO2-; R6 = H, alkyl; the dotted line represents either a double bond or a single bond] were prepared Reaction of 7β-[(Z)-2-(2-aminothiazol-4-yl)-2-[(8-hydroxy-2-oxo-1H-quinoline-5-yl) (carboxyl)methyloxyimino]acetamido]cephalosporanic acid di-Na salt with pyridine in the presence of NaI gave cephem (Z)-II isolated as α and β isomers. The title compds. in vitro exhibited MIC values of 0.1-0.78 μg/mL (against Staphylococcus aureus 209P JC-1) and MIC values of 0.78-1.56 μg/mL against Pseudomonas aeruginosa Number 12.

ΙI

146992-49-6P
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)
 (preparation of, as antibacterial agent)

RN 146992-49-6 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2-amino-4-thiazolyl)[[carboxy(1,2-dihydro-8-hydroxy-2-oxo-5-quinolinyl)methoxy]imino]acetyl]amino]-3-ethenyl-8-oxo-, disodium salt, [6R-( $6\alpha$ ,7 $\beta$ )]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.
Double bond geometry unknown.

### •2 Na

L9 ANSWER 34 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1993:212755 CAPLUS

DOCUMENT NUMBER: 118:212755

TITLE: Preparation of cephalosporin compounds

INVENTOR(S): Takamura, Norio; Saito, Kunio; Matsushita, Tadahiro;

Yamaguchi, Totaro

PATENT ASSIGNEE(S): Tanabe Seiyaku Co., Ltd., Japan

SOURCE: 'Jpn. Kokai Tokkyo Koho, 31 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 04261182	<b>A</b> 2	19920917	JP 1991-287408	19910808
JP 06086461	B4	19941102		
CA 2057129	AA	19930606	CA 1991-2057129	19911205
EP 544958	A1	19930609	EP 1991-311373	19911206
R: AT, BE, CH,	DE, DK	, ES, FR, GB	, GR, IT, LI, LU,	NL, SE
CN 1073444	Α	19930623	CN 1991-111604	19911218
PRIORITY APPLN. INFO.:			JP 1990-212040	A1 19900809
OTHER SOURCE(S):	MARPAT	118:212755		
CT				

$$\begin{array}{c|c} & & & & & & \\ & & & & & \\ & & & & \\ & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & &$$

Cephalosporin compds. [I; R1 = NH2, etc.; R2 = OH, etc.; R2 = CO2H, etc.; R4 = H, alkyl, alkenyl, CH2R (wherein R = nucleophilic radical such as AcO, pyridino, quinolino, thiazolylthio, etc.); R5 = CO2H, etc.; R6 = H, etc.; dotted line = saturation or unsatn.], useful as broad-spectrum antibacterial agents, are prepared A solution of DMF and POCl3 in CH2Cl2 was stirred at room temperature under Ar, cooled to -55° to -50°, treated with 13 g acid II (preparation given) in CH2Cl2 at -60° to -50°, and the solution was then treated with a suspension of MeC(OSiMe3):NSiMe3 and 5.43 g (syn)-I [R1 = Ph3CNH, R2 = 8-Ph2CHO, R3 = Ph2CHO2C, R4 = AcOCH2, R5 = CO2H, R6 = H, unsatd.]. The preferred dose was 5-40 mg/kg-day.

Ι

ΙI

IT 146992-49-6P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation of, as bactericide)

RN 146992-49-6 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2-amino-4-thiazolyl)[[carboxy(1,2-dihydro-8-hydroxy-2-oxo-5-quinolinyl)methoxy]imino]acetyl]amino]-3-ethenyl-8-oxo-, disodium salt, [6R-( $6\alpha$ ,7 $\beta$ )]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.
Double bond geometry unknown.

$$HO$$
 $O$ 
 $N$ 
 $H$ 
 $R$ 
 $R$ 
 $R$ 
 $CO_2H$ 
 $CO_2H$ 
 $CO_2H$ 

●2 Na

L9 ANSWER 35 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1993:143253 CAPLUS

DOCUMENT NUMBER: 118:143253

TITLE: In vitro susceptibilities of Actinobacillus

actinomycetemcomitans to a number of antimicrobial

combinations

AUTHOR(S): Pavicic, M. J. A. M. P.; Winkelhoff, A. J.; De Graaff,

J.

CORPORATE SOURCE: Dep. Oral Microbiol., Acad. Cent. Den., Amsterdam,

1081 BT, Neth.

SOURCE: Antimicrobial Agents and Chemotherapy (1992), 36(12),

2634-8

CODEN: AMACCQ; ISSN: 0066-4804

DOCUMENT TYPE: Journal LANGUAGE: English

AB The in vitro susceptibilities of A. actinomycetemcomitans to 14 antimicrobial combinations were studied by using the checkerboard titration technique. The results, expressed as the range of the fractional inhibitory concentration indexes, were as follows: for metronidazole or its hydroxymetabolite combined with cefixime, 0.2 to 0.6; for moxalactam, 0.2 to 0.6; for penicillin G, 0.3 to 0.6; for tobramycin, 0.8 to 2.0; for erythromycin, 0.8 to 1.7; for ciprofloxacin, 0.2 to 0.6; for tetracycline, 0.8 to 1.2. These observations indicated that the β-lactam antibiotics as well as ciprofloxacin act synergistically with both metronidazole and its hydroxymetabolite against A. actinomycetemocmitans. Synergistic interactions were independent of the individual MICs of the antibiotics tested. Erythromycin, tobramycin, and tetracycline combined with either metronidazole or its hydroxymetabolite showed additive to indifferent effects against the five strains of A. actinomycetemcomitans, with the fractional inhibitory concentration indexes ranging from 0.8 to 2.0.

A. actinomycetemcomitans was highly susceptible to ciprofloxacin (MIC of ciprofloxacin for 90% of strains tested, 0.010  $\mu$ g/mL) and cefixime (MIC of cefixime for 90% of strains tested, 0.8  $\mu$ g/mL). The results indicate that in patients who are allergic to penicillin, cefixime and ciprofloxacin may be useful alternative antibiotics in combination with metronidzole for the treatment of A. actinomycetemcomitans-associated

periodontitis.

IT 146505-62-6 146505-69-3

RL: BIOL (Biological study)

(Actinomyces actinomycetemcomitans sensitivity to)

RN 146505-62-6 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,
7-[[(2-amino-4-thiazolyl)[(carboxymethoxy)imino]acetyl]amino]-3-ethenyl-8oxo-, [6R-[6α,7β(Z)]]-, mixt. with 2-methyl-5-nitro-1Himidazole-1-ethanol (9CI) (CA INDEX NAME)

CM 1

CRN '79350-37-1

CMF C16 H15 N5 O7 S2

Absolute stereochemistry.

Double bond geometry as shown.

CM 2

CRN 443-48-1 CMF C6 H9 N3 O3

$$N$$
 $N$ 
 $N$ 
 $CH_2-CH_2-OH$ 

RN 146505-69-3 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,
7-[[(2-amino-4-thiazolyl) [(carboxymethoxy)imino]acetyl]amino]-3-ethenyl-8oxo-, [6R-[6α,7β(Z)]]-, mixt. with 2-(hydroxymethyl)-5-nitro-1Himidazole-1-ethanol (9CI) (CA INDEX NAME)

CM 1

CRN 79350-37-1

CMF C16 H15 N5 O7 S2

Absolute stereochemistry. Double bond geometry as shown.

2 CM

CRN 4812-40-2 CMF C6 H9 N3 O4

$$_{\text{O}_{2}\text{N}}^{\text{N}}$$
  $_{\text{CH}_{2}-\text{CH}_{2}-\text{OH}}^{\text{N}}$ 

L9 ANSWER 36 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1992:456006 CAPLUS

DOCUMENT NUMBER: TITLE:

117:56006

Direct compression method for cephalosporanic acid derivative tablets

INVENTOR(S):

Laly, Jean Louis; Lombardi, Roberto

PATENT ASSIGNEE(S):

Rhone-Poulenc Rorer S.A., Fr.

PCT Int. Appl., 13 pp.

SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

French

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9208463	A1	19920529	WO 1991-FR872	19911108
W: CA, JP, KR,	US			
RW: AT, BE, CH,	DE, DK	, ES, FR, GB	, GR, IT, LU, NL, SE	
FR 2669221	A1	19920522	FR 1990-14210	19901115
FR 2669221	B1	19930115		
CA 2094122	AA	19920516	CA 1991-2094122	19911108
CA 2094122	C	20040720		
EP 557389	A1	19930901	EP 1991-920476	19911108
EP 557389	B1	19940921		

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE JP 06502420 T219940317 JP 1992-500409 19911108 JP 3253074 B2 20020204 **T**3 ES 1991-920476 19911108 ES 2060417 19941116 US 1993-64049 19930514 US 5514383 Α 19960507 PRIORITY APPLN. INFO .: FR 1990-14210 19901115 WO 1991-FR872 19911108

AB Title tablets are prepared from mixts. containing 20-90% 7acylaminocephalosporanic acid derivs. and the balance excipients (CaCO3, CaSO4, starch, mannitol, fructose, etc.). A mixture of cefixime-3H2O 184.60, pregellified starch 48.98, CaHPO4, 2H2O 122.44, Mg stearate 2.03, and Avicel pH 102 is suitable for tabletting by direct compression.

125110-14-7, Cefixime trihydrate IT

RL: BIOL (Biological study)

(tabletting of, by direct compression)

125110-14-7 CAPLUS RN

5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, CN

7-[[(2Z)-(2-amino-4-thiazolyl)[(carboxymethoxy)imino]acetyl]amino]-3ethenyl-8-oxo-, trihydrate, (6R,7R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

H<sub>2</sub>O

L9 ANSWER 37 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1991:178375 CAPLUS

DOCUMENT NUMBER: 114:178375

Synergistic bactericidal compositions comprising TITLE:

decaplanin and cephalosporin derivatives

Seibert, Gerhard; Isert, Dieter; Klesel, Norbert

INVENTOR (S):

PATENT ASSIGNEE(S): Hoechst A.-G., Germany

SOURCE: Ger. Offen., 11 pp. CODEN: GWXXBX

DOCUMENT TYPE: Patent

LANGUAGE: German FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

DE 3909056	A1	19900920	DE 1989-3909056	19890318
EP 388510	A1	19900926	EP 1989-115520	19890823
R: AT, BE, CH,	DE, ES	, FR, GB,	GR, IT, LI, LU, NL, SE	
ZA 8906503	A	19900530	ZA 1989-6503	19890825
DK 8904208	Α	19900919	DK.1989-4208	19890825
AU 8940238	A1	19900920	AU 1989-40238	19890825
AU 625559	B2	19920716		
JP 02273624	A2	19901108	JP 1989-217617	19890825
HU 53539	A2	19901128	HU 1989-4415	19890825
HU 208087	В	19930830		
PRIORITY APPLN. INFO.:			DE 1989-3909056 A	19890318
OTHER SOURCE(S):	MARPAT	114:17837	75	
GI				

- \* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY AVAILABLE VIA OFFLINE PRINT \*
- AB Synergistic compns. useful for the prevention and treatment of bacterial inflammatory diseases comprise decaplanin (I) or I salt and a known cephalosporin antibiotic (Markush given). The compns. are especially useful against methicillin-resistant Staphylococcus, as shown by in-vitro studies on clin. isolates, using I-Cefpirome mixts.
- IT 133023-31-1
  - RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BIOL (Biological study) (bactericide, synergistic)
- RN 133023-31-1 CAPLUS
- CN Vancomycin, 22-0-(3-amino-2,3,6-trideoxy-3-C-methyl- $\alpha$ -L-arabino-hexopyranosyl)-2'-0-de(3-amino-2,3,6-trideoxy-3-C-methyl- $\alpha$ -L-lyxo-hexopyranosyl)-19-dechloro-2'-0-(6-deoxy- $\alpha$ -L-mannopyranosyl)-, mixt. with [6R-[6 $\alpha$ ,7 $\beta$ (Z)]]-7-[[(2-amino-4-thiazolyl)[(carboxymethoxy)imino]acetyl]amino]-3-ethenyl-8-oxo-5-thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid (9CI) (CA INDEX NAME)

CM 1

CRN 128441-18-9

CMF C72 H86 Cl N9 O28

CM 2

CRN 79350-37-1 CMF C16 H15 N5 O7 S2

Absolute stereochemistry.
Double bond geometry as shown.

L9 ANSWER 38 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1991:150303 CAPLUS

DOCUMENT NUMBER: 114:150303

TITLE: Determination of water in drug substances by Karl

Fischer method with water vaporizer

AUTHOR(S): Kitagawa, Teruyuki; Hara, Mitsue; Yokobayashi,

Shizuka; Kawabata, Tetsuo; Koda, Shigetaka; Yasuda,

Tsutomu

CORPORATE SOURCE: Anal. Res. Lab., Fujisawa Pharm. Co., Ltd., Osaka,

532, Japan

SOURCE: Bunseki Kagaku (1991), 40(1), T9-T13

CODEN: BNSKAK; ISSN: 0525-1931

DOCUMENT TYPE: Journal LANGUAGE: Japanese

AB The accuracy and anal. precision of the Karl Fishcer (KF) method with water vaporizer was enough high compared with those of the direct KF method and this method is applied to pharmaceuticals which interfere with KF reagents. Suitable temperature for vaporizing H2O was 150°, but in some cases, 10° below the decomposition point was appropriate. The effects of desiccants for a carrier gas and the heating temperature upon the blank value were examined It was found that the volume of KF reagent consumed for a sample titration must be corrected using the blank value obtained in the same titration time in all cases.

IT 125110-14-7, Cefixime trihydrate

RL: AMX (Analytical matrix); ANST (Analytical study)

(water determination in, by Karl Fischer method, with water vaporizer)

RN 125110-14-7 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,

7-[[(2Z)-(2-amino-4-thiazolyl)[(carboxymethoxy)imino]acetyl]amino]-3-

ethenyl-8-oxo-, trihydrate, (6R,7R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

●3 H<sub>2</sub>O

L9 ANSWER 39 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1990:429182 CAPLUS

DOCUMENT NUMBER: 113:29182

TITLE: Dehydration effect on the stability of cefixime

triĥydrate

AUTHOR(S): Kitamura, Satoshi; Koda, Shigetaka; Miyamae, Akira;

Yasuda, Tsutomu; Morimoto, Yukiyoshi

CORPORATE SOURCE: Anal. Res. Lab., Fujisawa Pharm. Co., Ltd., Osaka,

532, Japan

SOURCE: International Journal of Pharmaceutics (1990), 59(3),

217-24

CODEN: IJPHDE; ISSN: 0378-5173

DOCUMENT TYPE: Journal LANGUAGE: English

AB Partially dehydrated cefixime trihydrate was unstable due to a highly disordered crystal structure caused by loss of its water of crystallization. It was also confirmed that cefixime trihydrate stored at a relative humidity below its critical value was less stable than the trihydrate stored under moist conditions. On the other hand, completely dehydrated cefixime trihydrate was relatively stable since it underwent transformation to a new anhydrous crystal form which did not contain water capable of participating in the hydrolytic reaction. It was suggested that the degradation mechanism under conditions of dryness differed from that under conditions of humidity, since not only the appearance but also the particular species of degradation products were completely different under the two sets of conditions.

IT 125110-14-7, Cefixime trihydrate

RL: PRP (Properties)

(stability of, dehydration effect on)

RN 125110-14-7 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,

7-[[(2Z)-(2-amino-4-thiazolyl)[(carboxymethoxy)imino]acetyl]amino]-3-ethenyl-8-oxo-, trihydrate, (6R,7R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

$$HO_2C$$
 $O$ 
 $N$ 
 $Z$ 
 $N$ 
 $NH_2$ 
 $HN$ 
 $R$ 
 $R$ 
 $R$ 
 $CH_2$ 

# ●3 H<sub>2</sub>O

L9 ANSWER 40 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1990:62490 CAPLUS

DOCUMENT NUMBER: 112:62490

TITLE: Effect of grinding on the solid-state stability of

cefixime trihydrate

AUTHOR(S): Kitamura, Satoshi; Miyamae, Akira; Koda, Shigetaka;

Morimoto, Yukiyoshi

CORPORATE SOURCE: Anal. Res. Lab., Fujisawa Pharm. Co., Ltd., Osaka,

532, Japan

SOURCE: International Journal of Pharmaceutics (1989), 56(2),

125-34

CODEN: IJPHDE; ISSN: 0378-5173

DOCUMENT TYPE: Journal LANGUAGE: English

AB The effect of grinding on the physicochem. properties of cefixime trihydrate (I) was studied by means of x-ray diffraction anal., SEM, DSC equilibrium water amts. and color difference measurement  $(\Delta E)$ . Crystalline I was confirmed to change to a non-crystalline solid after 4 h of grinding in a ball mill, since x-ray diffraction peak intensities decreased with

increasing grinding time. Dehydration temperature of ground I also lowered

with

increasing grinding time, and the activation energy for dehydration of intact I and the samples ground 4 h (amorphous form) were calculated by Kissinger's method to be 72.4 kcal/mol and 67.5 kcal/mol, resp. The decreased crystallinity with grinding is presumably due to an increase of water mols. having greater freedom of movement in the crystal lattice. The overall decomposition of solid-state I could be expressed by pseudo first-order reaction, and the crystallinity of the ground sample was estimated by an equation expressing the overall decomposition rate constant; which is the sum of the decomposition in 100% crystalline and in 0% crystalline (amorphous)

Kinetic studies of discoloration of ground I showed an increase in the apparent rate constant for discoloration with the increase in the grinding time.

IT 125110-14-7

RL: PRP (Properties)

(stability of, in solid state, grinding effect on)

RN 125110-14-7 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2Z)-(2-amino-4-thiazolyl)[(carboxymethoxy)imino]acetyl]amino]-3-ethenyl-8-oxo-, trihydrate, (6R,7R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

●3 H<sub>2</sub>O

L9 ANSWER 41 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1990:25681 CAPLUS

DOCUMENT NUMBER: 112:25681

TITLE: Antiulcer agents containing cefixime (salts)

INVENTOR(S): Ono, Takaharu; Tomoi, Masaaki

PATENT ASSIGNEE(S): Fujisawa Pharmaceutical Co., Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 4 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PATENT NO.	KIND	DATE	APPLICATION :	NO.	DATE
	JP 01149728	A2	19890612	JP 1987-3082	14	19871204
PRIC	RITY APPLN. INFO.:			JP 1987-3082	14	19871204
AB	Antiulcer agents co	ntain c	efixime (I)	or its salts.	I at 320	0 mg/kg p.o.
	showed 90.4% inhibi	tion ag	ainst stress	s-induced ulce	r in rats.	LD50 of I
	was ≥10,000 mg/kg p	.o. in	rats. Table	ets were formu	lated conta	aining I
	93.5, CMC Ca 3.7, M	g stear	ate 1.9, and	d silica 0.9 w	eight%.	
IT	124506-28-1		•			
	RL: BIOL (Biologica	l study	) ′			
	(antiulcer agent	s conta	ining)			
RN	124506-28-1 CAPLUS					
CN	5-Thia-1-azabicyclo	[4.2.0]	oct-2-ene-2-	-carboxylic ac	id,	
	7-[[(2-amino-4-thia	zolyl)[	(carboxymeth	noxy)imino]ace	tyl]amino]	-3-ethenyl-8-
	oxo-, disodium salt	, [6R-[	6α, 7α(Z)]]-	(9CI) (CA IN	DEX NAME)	

Absolute stereochemistry.

Double bond geometry as shown.

### ●2 Na

L9 ANSWER 42 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1989:423290 CAPLUS

DOCUMENT NUMBER:

111:23290

TITLE:

Preparation of thiadiazolyl(aminoethoxyimino)acetamido

cephalosporin compounds as antibacterial agents

INVENTOR(S):

Nishizawa, Susumu; Muro, Hiroyuki; Kasai, Masayasu; Hatano, Satoru; Kamiya, Syouzi; Kakeya, Nobuharu;

Kitao, Kazuhiko

PATENT ASSIGNEE(S):

Kyoto Pharmaceutical Industries, Ltd., Japan

SOURCE:

Eur. Pat. Appl., 35 pp. CODEN: EPXXDW

DOCUMENT TYPE:

: Patent English

LANGUAGE:
FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
EP 293771	<b>A</b> 2	19881207	EP 1988-108456		19880527
EP 293771	<b>A</b> 3	19901017			
R: AT, BE, CH	, DE, ES	, FR, GB,	IT, LI, NL, SE		
JP 01056682	A2	19890303	JP 1988-98539		19880420
AU 8816309	A1	19881201	· AU 1988-16309		19880517
DK 8802871	Α	19881201	DK 1988-2871		19880525
US 4943567	Α	19900724	US 1988-228714		19880527
AU 9063814	A1	19910228	AU 1990-63814		19901003
AU 628664	B2	19920917			
PRIORITY APPLN. INFO.:		•	JP 1987-136647	Α	19870530
OTHER SOURCE(S):	CASREA	CT 111:232	290; MARPAT 111:23290		
GT					

Cephalosporin derivs. (I; R1, R5 = H, protecting group; R2 = alkyl, cycloalkyl; R3 = H, alkenyl, acyloxymethyl, carbamoyloxymethyl, heterocyclylthiomethyl, etc.; R4 = H, ester residue; X = CH, N) and their pharmacol. acceptable salts are prepared A mixture of syn-II, III, pyridine, and POCl3 in CH2Cl2 was stirred at -12° to -15° to give syn-I (R1 = CO2CMe3, R2 = Et, R3 = 1,3,4-thiadiazol-2-ylthiomethyl, R4 = Ph2CH, R5 = HCO, X = CH), which was hydrolyzed to give the acid syn-I (R4 = H, others remain unchanged) (IV). Deprotection of IV with concentrated HCl

I.

II

MeOH gave syn-I·2HCl (R1 = R4 = R5 = H, others = same), which showed MIC of 0.39  $\mu$ g/mL against Staphylococcus aureus. A parenteral solution was made from 1 g syn-I·2HCl (R1 = R4 = R5 = H, R2 = Me, R3 = CH2OAc, X = CH) and 135 mg Na2CO3 in 20 mL distilled H2O.

IT 121102-80-5P

in

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation of, as antibacterial agent)

RN 121102-80-5 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,
7-[[[[1-(aminomethyl)propoxy]imino](2-amino-4-thiazolyl)acetyl]amino]-3ethenyl-8-oxo-, dihydrochloride, [6R-[6α,7β(Z)]]- (9CI) (CA
INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

# ●2 HCl

L9 ANSWER 43 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1988:549222 CAPLUS

DOCUMENT NUMBER: 109:149222

TITLE: Preparation of cephalosporin derivatives

INVENTOR(S): Nakagawa, Susumu; Fukatsu, Hiroshi; Murase, Satoshi

PATENT ASSIGNEE(S): Banyu Pharmaceutical Co., Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 6 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent

LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE	
JP 63119488	A2	19880524	JP 1986-261844	19861105	
PRIORITY APPLN. INFO.:			JP 1986-261844	19861105	
OTHER SOURCE(S):	MARPAT	109:149222			
GI					

$$\begin{array}{c|c}
N & CCONH \\
\parallel & NOC (CO_2H) = CH_2 \\
\hline
 & CO_2H \\
\hline
 & CO_2H$$

AB Title derivs. I [R1 = H, NH2; R2 = H, halo, (substituted) lower alkyl, lower alkenyl, lower alkoxy, or alkylthio], their nontoxic salts, or physiol. hydrolyzable nontoxic esters are prepared (Z)-2-(1-tert-Butoxycarbonylvinyloxyimino)-2-(2-tritylaminothiazol-4-yl)acetic acid (preparation given) was stirred with POCl3 and DMF in THF at 0° for 1 h then treated with a solution containing p-methoxybenzyl 7-amino-3-(methylthio)-3-

cephem-4-carboxylate.HCl and N,O-bis(trimethylsilyl)acetamide in AcOEt at 0° for 1 h to give corresponding p-methoxybenzyl acetamidocephemcarboxylate derivative, which was deprotected by treating with CF3CO2H and anisole at room temperature for 1 h to give 31.6% I (R1 = NH2, R2 = SMe) (II). II in vitro exhibited MIC value of 0.2  $\mu$ g/mL against Escherichia coli NIHJ JC2.

IT 116797-41-2P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation of, as antibiotic)

RN 116797-41-2 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2-amino-4-thiazolyl)[[(1-carboxyethenyl)oxy]imino]acetyl]amino]-3-ethenyl-8-oxo-, disodium salt, [6R-[6 $\alpha$ ,7 $\beta$ (Z)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

2 Na

9 ANSWER 44 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1988:406314 CAPLUS

DOCUMENT NUMBER:

109:6314

TITLE:

Preparation of [(pyridonylmethoxyimino)acetamido]cephe

mcarboxylic acid derivatives as antibiotics

INVENTOR(S):

Zama, Yoshiyuki; Ishiyama, Nobuo; Saita, Tsuneo; Naito, Takanobu; Hirose, Masao; Yokoyama, Masaaki; Asano, Taiji; Senda, Hisato; Sekine, Keiji; Sanai,

Shigeru

PATENT ASSIGNEE(S):

Kaken Pharmaceutical Co., Ltd., Japan

SOURCE:

Eur. Pat. Appl., 41 pp.

DOCUMENT TYPE:

CODEN: EPXXDW

DOCUMENT TIPE

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.

KIND DATE

APPLICATION NO.

DATE

											~		
. EP	251299			A2	19	880	107		ΕP	1987-109416			19870630
EP	251299			<b>A</b> 3	19	891	011						
EP	251299			B1	. 19	940	831						
	R: A.	Г, ВЕ,	CH,	DE,	ES, E	R,	GB,	GR,	rı	C, LI, LU, NL,	SE		
US	4822786	5		Α	19					1987-63077			19870617
CA	1283404	1		A1	19	910	423		CA	1987-539868			19870617
HU	44259			A2	19	9880	229		HU	1987-2882			19870625
HU	198500			В	19	891	030						
HU	56099			A2 B A2	19	910	729		HU	1989-2782			19870625
HU	209126			В	19	9940	328						
HU	56071			A2		910	729		HU	1990-8257			19870625
HU	207296				19	930	329						
AU	877493	1				9880	107		ΑU	1987-74931			19870629
AU	597676			B2		900	607						
CN	8710459	90		Α	19	9880	113		CN	1987-104590			19870630
CN	1022036	5		В	19	930	908						
. ES	2062974	4 .		Т3	19		101		ES	1987-109416			19870630
JP	6314688	87		A2	19	9880	618		JΡ	1987-162296			19870701
JP	0603126			B4	19	940	427						
JP	6315238					9880	624		JP	1987-203494			19870818
JP	0605170	06		B4	19	9940	706						
US	4883879	9		Α	19	9891	128		US	1989-296765			19890113
JP	0228888	84		A2		9901	128		JP	1989-341529			19891229
JP	060864	52		B4	19	9941	102						
CA	1333713	3		A1	19	9941	.227			1990-615847			19900823
AU	9062143					9901	220		AU	1990-62143			19900904
AU	627067			B2		920	813						
AU	9219628	В		A1		920	910		ΑU	1992-19628			19920710
AU	635174			B2	19	930	311						
PRIORIT	Y APPLN	. INFO	.:						JP	1986-152706	A		19860701
									JP	1986-191590	Α		19860818
									CA	1987-539868	A.	3	19870617
										1987-63077	A:	3	19870617
OTHER S	OURCE(S)	) :		CASI	REACT	109	:631	L4;	MAF	RPAT 109:6314			

OTHER SOURCE(S)

$$R^{2}HN$$
 $S$ 
 $N$ 
 $CCONH$ 
 $S$ 
 $CH_{2}X$ 
 $CO_{2}R^{5}$ 
 $NO^{4}$ 
 $NO^{4}$ 
 $OR^{3}$ 
 $IV$ 

AB The title compds. I [R1 = H, halo, MeO, (substituted) vinyl, CH2A wherein A = H, N3, acyloxy, carbamoyloxy, (substituted) heterocyclyl, heterocyclylthio], useful as antibiotics, were prepared from II (R2 = H, amino-protecting group; R3, R4 = H, OH-protecting group), III (X = Cl, Br, iodine, acetoxy; R5 = H, CO2H-protecting group), and IV. Reaction of p-methoxybenzyl (6R,7R)-7-[(Z)-2-(2-tritylaminothiazol-4-yl)-2-(1,5-dibenzhydryloxy-4-pyridon-2-ylmethoxyimino)acetamido]-3-chloromethyl-3-cephem-4-carboxylate with Na 1,2,3-thiadiazol-5-thiolate, followed by deprotection and workup, gave (6R,7R)(Z)-I (R1 = 1,2,3-thiadiazol-5-ylthiomethyl) Na salt (V). V in vitro exhibited a MIC of 6.25 μg/mL against Staphylococcus aureus FDA 209-P.

IT 114830-52-3P 114904-05-1P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation of, as antibiotic)

RN 114830-52-3 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2-amino-4-thiazolyl)[[(1,4-dihydro-1,5-dihydroxy-4-oxo-2-pyridinyl)methoxy]imino]acetyl]amino]-3-ethenyl-8-oxo-, monosodium salt, [6R-[ $6\alpha$ ,7 $\beta$ (Z)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

Na

RN 114904-05-1 CAPLUS CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2-amino-4-thiazolyl)[[(1,4-dihydro-1,5-dihydroxy-4-oxo-2-pyridinyl)methoxy]imino]acetyl]amino]-3-ethenyl-8-oxo-, [6R-[6 $\alpha$ ,7 $\beta$ (Z)]]-, mono(trifluoroacetate) (salt) (9CI) (CA INDEX NAME)

CM 1

CRN 114876-06-1 CMF C20 H18 N6 O8 S2

Absolute stereochemistry.

Double bond geometry as shown.

CM 2

CRN 76-05-1 CMF C2 H F3 O2

L9 ANSWER 45 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1986:33941 CAPLUS

DOCUMENT NUMBER: 104:33941

TITLE: Cephem derivatives

PATENT ASSIGNEE(S): Meiji Seika Kaisha, Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 24 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 60105683	A2	19850611	JP 1983-212461	19831114
JP 02027998	B4	19900620		·
PRIORITY APPLN. INFO.:			JP 1983-212461	19831114
CT ·				

Ι

$$\begin{array}{c|c}
N & C - CONH & S \\
0 & N & R3
\end{array}$$

$$\begin{array}{c|c}
0 - CZR^2 & R4 \\
0 & O
\end{array}$$

$$\begin{array}{c|c} N & C - CO_2K \\ \hline Ph_3CNH & S & O - C \\ \hline & O - C \\ & & O \end{array}$$

AB Cephem derivs. (I; R1 = NH2, protected NH2; R2, R4 = CO2H, protected CO2H; R3 = H, halo, alkylthio, etc.; Z = C2-10 alkylene, phenylene, cycloalkylene), effective antibacterials at 0.025-12.5 μg/mL were prepared Thus, 5% HCl was added to a suspension of 380 mg syn-II in EtoAc-THF to pH 2.5 under cooling, 70 mg 1-hydroxybenzotriazole and 250 mg III were added to solution, 103 mg DCC added to 5° and stirred to give 310 mg syn-I (R1 = Ph3CNH, R2Z = p-C6H4CO2CHPh2, R3 = 1-methyl-1,2,3,4-

tetrazol-5-ylthiomethyl, R4 = CO2CHPh2). IT 99743-93-8P 99744-01-1P RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation) (preparation and antibacterial activity of) 99743-93-8 CAPLUS RN5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, CN 7-[[(2-amino-4-thiazolyl)[(3-carboxy-1-oxopropoxy)imino]acetyl]amino]-3ethenyl-8-oxo-,  $[6R-[6\alpha,7\beta(Z)]]$ -, mono(trifluoroacetate) (9CI) (CA INDEX NAME) CM 1 CRN 99743-92-7 CMF C18 H17 N5 O8 S2

Absolute stereochemistry. Double bond geometry as shown.

CRN 76-05-1 CMF C2 H F3 O2

Absolute stereochemistry. Double bond geometry as shown.

CM 2

CRN 76-05-1 CMF C2 H F3 O2

ANSWER 46 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN L9

ACCESSION NUMBER:

1985:148993 CAPLUS

DOCUMENT NUMBER:

102:148993

TITLE:

3-phosphonium and 3-phosphoranylidenecephems

INVENTOR(S):

Takaya, Takao; Takasugi, Hisashi; Masugi, Takashi;

Yamanaka, Hideaki; Kawabata, Kohji

PATENT ASSIGNEE(S):

Fujisawa Pharmaceutical Co., Ltd., Japan

SOURCE:

U.S., 82 pp. Cont.-in-part of U.S. 4,409,214.

CODEN: USXXAM

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

9

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 4487927	A	19841211	US 1982-341621	19820122
US 4409214 .	A	19831011	US 1980-205334	19801110
ZA 8006977	Α	19811028	ZA 1980-6977	19801111
AT 37028	E	19880915	AT 1984-100915	19801115
AT 86987	E	19930415	AT 1987-104893	19801115
US 4423213	Α	19831227	US 1981-261618 .	19810507
ES 507973 .	A1	19821001	ES 1981-507973	19811215
JP 58135894	A2	19830812	JP 1983-9235	19830121
JP 05001271	B4 .	19930107		

US 4559334	A	19851217	US	1983-543880		19831020
US 4904652	Α	19900227	US	1985-785048		19851009
US 4731443	Α.	19880315	US	1986-889189		19860724
SU 1508962	A3	19890915	SU	1987-4202592		19870519
US 4960889	Α	19901002	US	1990-462347		19900103
US 5026695	Α	19910625	US	1990-461340		19900105
JP 02223544	A2	19900905	JP	1990-11048		19900119
JP 06078290	B4	19941005				
US 5110921	Α	19920505	US	1990-583304		19900917
US 5594132	Α	19970114	US	1991-684194		19910412
US 5252731	A	19931012	US	1992-831504		19920205
PRIORITY APPLN. INFO.:			GB	1979-39985	Α	19791119
			GB	1980-4335	Α	19800208
			GB	1980-12991	Α	19800421
			GB	1980-22920	Α	19800714
			US	1980-205334	A2	19801110
			US	1981-261618	A2	19810507
			US	1980-206831	A3	19801114
		•	EP	1984-100915	Α	19801115
			EP	1987-104893	Α	19801115
			US	1982-341621	Α	19820122
			US	1982-428970	A2	19820930
			US	1983-489236	В1	19830428
			GB	1983-23034	Α	19830826
•			US	1984-653041	АЗ	19840921
			US	1985-785048	Α3	19851009
			US	1986-889189	B3	19860724
	•		US	1987-127929		19871202
				1990-462347		19900103
				1990-461340		19900105
			US	1990-583304	Α3	19900917

GΙ

RON = CR1CONH

The title compds. I [R = alkyl, (un)esterified carboxyalkyl; R1 = (un)protected aminothiazol-4-yl; R2 = CH2P+R43X-, CH:PR43; R3 = H, protective group; R4 = aryl; X = halogen] were prepared as intermediates for 3-vinylcephems. Thus II was obtained by reaction of PPh3 and NaI with the corresponding 3-chloromethylcephem which was prepared from cephalosporin C in 3 steps.

TT 79350-11-1P 79350-44-0P 79350-82-6P 86027-36-3P 90467-43-9P 90467-53-1P

95759-13-0P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and esterification of)

RN 79350-11-1 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,
7-[[(2-amino-4-thiazolyl) (methoxyimino)acetyl]amino]-3-ethenyl-8-oxo-,
monohydrochloride, [6R-[6α,7β(Z)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

● HCl

RN 79350-44-0 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,
7-[[(2Z)-(2-amino-4-thiazolyl)(methoxyimino)acetyl]amino]-3-ethenyl-8-oxo, monosodium salt, (6R,7R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

RN 79350-82-6 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2-amino-4-thiazolyl)[(carboxymethoxy)imino]acetyl]amino]-3-ethenyl-8-oxo-, disodium salt, [6R-[6 $\alpha$ ,7 $\beta$ (Z)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

●2 Na

RN 86027-36-3 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2-amino-4-thiazolyl)[[2-(diphenylmethoxy)-2-oxoethoxy]imino]acetyl]amino]-3-ethenyl-8-oxo-, monosodium salt, [6R-[6 $\alpha$ ,7 $\beta$ (Z)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

Na

RN 90467-43-9 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2-amino-4-thiazolyl)[(2-propynyloxy)imino]acetyl]amino]-3-ethenyl-8-oxo-, monosodium salt, [6R-[6 $\alpha$ ,7 $\beta$ (Z)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.
Double bond geometry as shown.

Na

RN 90467-53-1 CAPLUS CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2-amino-4-thiazolyl) (ethoxyimino)acetyl]amino]-3-ethenyl-8-oxo-, monosodium salt, [6R-[6 $\alpha$ ,7 $\beta$ (Z)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

Na

RN 95759-13-0 CAPLUS
CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,
7-[[(2-amino-4-thiazolyl)[[2-[(4-nitrophenyl)methoxy]-2-oxoethoxy]imino]acetyl]amino]-3-ethenyl-8-oxo-, monosodium salt,

$$[6R-[6\alpha,7\beta(Z)]]$$
 - (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

Na

IT 88621-04-9P

RN 88621-04-9 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,
7-[[(2-amino-4-thiazolyl)[[2-[(4-nitrophenyl)methoxy]-2oxoethoxy]imino]acetyl]amino]-3-ethenyl-8-oxo-, monohydrochloride,
[6R-[6α,7β(Z)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

● HCl

## Page 101

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation, esterification, and bactericidal activity of)

RN 79369-28-1 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,

7-[[(2-amino-4-thiazolyl)[(carboxymethoxy)imino]acetyl]amino]-3-ethenyl-8-oxo-, monohydrochloride, [6R-[6 $\alpha$ ,7 $\beta$ (Z)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

● HCl

RN 90467-54-2 CAPLUS

CN Pyridinium, 2-[[[1-(2-amino-4-thiazolyl)-2-[(2-carboxy-3-ethenyl-8-oxo-5-thia-1-azabicyclo[4.2.0]oct-2-en-7-yl)amino]-2oxoethylidene]amino]oxy]methyl]-1-methyl-, chloride, [6R[6α,7β(Z)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.
Double bond geometry as shown.

• cl -

90467-55-3 CAPLUS RN

Pyridinium, 3-[[[[1-(2-amino-4-thiazoly1)-2-[(2-carboxy-3-etheny1-8-oxo-5-CNthia-1-azabicyclo[4.2.0]oct-2-en-7-yl)amino]-2oxoethylidene]amino]oxy]methyl]-1-methyl-, chloride, [6R- $[6\alpha, 7\beta(Z)]$  - (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

● Cl -

CAPLUS COPYRIGHT 2006 ACS on STN ANSWER 47 OF 52 L9

1985:113176 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 102:113176

Novel cephem compounds TITLE:

PATENT ASSIGNEE(S): Meiji Seika Kaisha, Ltd., Japan Jpn. Kokai Tokkyo Koho, 18 pp. SOURCE:

CODEN: JKXXAF

#### Page 103

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE	
JP 59184186	A2	19841019	JP 1983-57465	19830401	
PRIORITY APPLN. INFO.:			JP 198357465	19830401	
GI .					

$$\begin{array}{c|c}
N & CCONH & S \\
\parallel & NO_2CR^1 & N \\
& & R^3
\end{array}$$

AB Cephems I (R = amino, protein amino; R1 = alkyl; R2 = vinyl, alkylthio, CH:CHCO2R4, CH2CO2R5; R3 = CO2H, protected carboxyl; R4; R5 = H, alkyl) were prepared Thus, amidation of syn-2-(2-tritylaminothiazol-4-yl)-2-(pivaloyloxyimino)acetic acid with diphenylmethyl 7-amino-3-vinyl-3-cephem-4-carboxylate followed by hydrolysis with Cl3CCO2H gave syn-I.Cl3CCO2H (R = NH2, R1 = Me3C, R2 = vinyl, R3 = CO2H). The latter compound showed broad spectrum bactericidal activity.

IT 94796-36-8P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation and bactericidal activity of)

Ι

RN 94796-36-8 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2-amino-4-thiazolyl)[(2,2-dimethyl-1-oxopropoxy)imino]acetyl]amino]-3-ethenyl-8-oxo-, [6R-[6 $\alpha$ ,7 $\beta$ (Z)]]-, trichloroacetate (9CI) (CA INDEX NAME)

CM 1

CRN 94796-35~7 CMF C19 H21 N5 O6 S2

Absolute stereochemistry.

Double bond geometry as shown.

CM 2

CRN 76-03-9 CMF C2 H Cl3 O2

L9 ANSWER 48 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1984:438270 CAPLUS

Correction of: 1982:181061

DOCUMENT NUMBER: 101:38270

Correction of: 96:181061

TITLE: 7-Acylamino-3-vinylcephálosporanic acid derivatives

INVENTOR(S): Takaya, Takao; Takasugi, Hisashi; Masugi, Takashi;

Yamanaka, Hideaki; Kawabata, Kohji

PATENT ASSIGNEE(S): Fujisawa Pharmaceutical Co., Ltd., Japan

SOURCE: Eur. Pat. Appl., 285 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 9

PATENT INFORMATION:

DAMENIM NO	WEND	D 3 (T) D	3 DD1 TG2 MTO11 310	D. 2. TT
PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 30630	A2	19810624	EP 1980-107075	19801115
EP 30630	A3	19810909		
EP 30630	B1	19870401		
R: AT, BE, CH,	DE, FR,	GB, IT, LU	, NL, SE	
ZA 8006977	A	19811028	ZA 1980-6977	19801111
CA 1235414	A1	19880419	CA 1980-364436	19801112
FI 8003558	Α	19810520	FI 1980-3558	19801113
FI 74970	В	19871231		
FI 74970	C	19880411		
EP 123024 ·	A2	19841031	EP 1984-100915	19801115

EP	123024			A3	19850313				
EP	123024			B1					
	R: AT,	BE,	CH,	DE,	FR, GB, IT,				
	26280			$\mathbf{E}$	19870415		1980-107075		19801115
	244637			<b>A1</b>	19871111	EP	1987-104893		19801115
EP	244637			B1	19930317				
		BE,	CH,	DE,	FR, GB, IT,				
	37028			E	19880915		1984-100915		19801115
	86987			E	19930415		1987-104893		19801115
	8064442			A1	19810528	AU	1980-64442		19801117
	543301			B2	19850418				
	8004917			A	19810619		1980-4917		19801118
	8003470			`A .		NO	1980-3470		19801118
	160921			В					
	160921			C					
	496948			A1	19820501		1980-496948		19801118
	1186087			A3	19851015		1980'-3009474		19801118
	56086187			A2	19810713	JP	1980-163989		19801119
	63020435			B4	19880427		1001 507070		10011015
	507972			A1	19821001		1981-507972		19811215
	507973			A1	19821001		1981-507973 1985-785048		19811215
	4904652			A	19900227				19851009
	62277391 03014832	•		A2 B4	19871202 19910227	UP	1987-44400		19870226
	1508962	•		A3	19890915	CII	1987-4202592		19870519
	63146863		•	A2	19880618		1987-4202532		19871117
	02025905			. B4	19900606	. 0P	1307-230253		190/111/
	63152387			A2	19880624	.TD	1987-290248		19871117
	03132307			B4	19910610	UF	1907-290240		190/111/
	63152388			A2	19880624	.TD	1987-290249		19871117
	03069353			B4	19911031	<u>.</u>	,		150/111/
	63152385			A2	19880624	дÞ	1987-290250		19871117
	03038277			B4	19910610	01	130, 130130		150,111,
	63152370			A2	19880624	JР	1987-290251		19871117
	03033712			B4	19910520				
	63152371			A2	19880624	JP	1987-290252		19871117
	02019828			B4	19900507				
US	4960889			Α	19901002	US	1990-462347		19900103
US	5026695			Α	19910625	US	1990-461340		19900105
US	5594132			Α	19970114	US	1991-684194		19910412
. Jb	06279452			A2	19941004	JP	1991-201550		19910510
JP	07010870			B4	19950208				•
PRIORITY	APPLN.	INFO.	:				1979-39985	Α	19791119
• .							1980-4335	Α	19800208
							1980-12991	A	19800421
	•						1980-22920	Α	19800714
		• .					1980-206831		19801114
							1980-107075	P	19801115
							1984-100915	A	19801115
	•				•		1987-104893	A	19801115
							1983-489236		19830428
							1985-785048		19851009
	•						1986-889189		19860724
							1987-127929		19871202
OMITTED CO	Amar (a)			07.05	TEN CITE 101 201		1990-461340	A3	19900105
GI GI	OURCE(S):			CASI	CEACI IUI:382	2/U; MA	ARPAT 101:38270		
GI.									

RXCONH 
$$S$$
  $CH = CH_2$   $CO_2R^1$   $I$ 

AB Vinylcephems I [R = (un)substituted aminoheterocyclic, R2SO2NHC6H4; R1 = H, protective group; R2 = alkyl; X = (un)substituted alkylene] were prepared Thus, I (R = 3-MeSO2NHC6H4, R1 = H, X = H2NCH, II) was obtained by acylating a 7-aminocephem with 3-MeSO2NHC6H4CH(NH2)CO2H.

7-Amino-3-vinyl-3-cephem-4-carboxylic acid was obtained from the hydroxymethylcephem via the chloromethyl derivative and the triphenylphosphonium iodide which was treated with CH2O. II had a min. inhibitory concentration against Staphylococcus aureus 209 P JC-1 of 1.56 μg/mL.

IT 79350-11-1P 79350-44-0P 79350-82-6P 90467-43-9P 90467-53-1P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and esterification of)

RN 79350-11-1 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2-amino-4-thiazolyl) (methoxyimino)acetyl]amino]-3-ethenyl-8-oxo-, monohydrochloride, [ $6R-[6\alpha,7\beta(Z)]$ ]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

#### ● HCl

RN 79350-44-0 CAPLUS
CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,
7-[[(2Z)-(2-amino-4-thiazolyl) (methoxyimino)acetyl]amino]-3-ethenyl-8-oxo, monosodium salt, (6R,7R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.
Double bond geometry as shown.

Page 107

Na

RN 79350-82-6 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2-amino-4-thiazolyl)[(carboxymethoxy)imino]acetyl]amino]-3-ethenyl-8-oxo-, disodium salt, [6R-[6 $\alpha$ ,7 $\beta$ (Z)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

$$HO_2C$$
 $O$ 
 $R$ 
 $R$ 
 $R$ 
 $R$ 
 $CH_2$ 

●2 N=

RN 90467-43-9 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,
7-[[(2-amino-4-thiazolyl)[(2-propynyloxy)imino]acetyl]amino]-3-ethenyl-8oxo-, monosodium salt, [6R-[6α,7β(Z)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

Na

RN 90467-53-1 CAPLUS
CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,
7-[[(2-amino-4-thiazolyl)(ethoxyimino)acetyl]amino]-3-ethenyl-8-oxo-,

monosodium salt,  $[6R-[6\alpha,7\beta(Z)]]-(9CI)$  (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

Na

Absolute stereochemistry.

Double bond geometry as shown.

● Cl -

RN 90467-55-3 CAPLUS
CN Pyridinium, 3-[[[[1-(2-amino-4-thiazolyl)-2-[(2-carboxy-3-ethenyl-8-oxo-5-thia-1-azabicyclo[4.2.0]oct-2-en-7-yl)amino]-2 oxoethylidene]amino]oxy]methyl]-1-methyl-, chloride, [6R-[6α,7β(Z)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

• cl -

 oxo-, monohydrochloride, [6R-[6 $\alpha$ ,7 $\beta$ (Z)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

## ● HCl

L9 ANSWER 49 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1984:406933 CAPLUS

DOCUMENT NUMBER: 101:6933

TITLE: 7-Acylamino-3-vinylcephalosporanic acid derivatives INVENTOR(S): Takaya, Takao; Takasugi, Hisashi; Masugi, Takashi;

Yamanaka, Hideaki; Kawabata, Kohji

PATENT ASSIGNEE(S): Fujisawa Pharmaceutical Co., Ltd. , Japan

SOURCE: U.S., 80 pp. Cont.-in-part of U.S. Ser. No. 205,334.

CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 9

PATENT INFORMATION:

PATENT N	o. K	CIND	DATE	API	PLICATION NO.	DATE
US 44232	13	A	19831227	US	1981-261618	19810507
US 44092	14	A	19831011	US	1980-205334	19801110
ZA 80069	77	A	19811028	zA	1980-6977	19801111
AT 37028		E	19880915	AT	1984-100915	19801115
AT 86987		E	19930415	AT	1987-104893	19801115
ES 50797	3	A1	19821001	ES	1981-507973	19811215
US 44879	27	A	19841211	US	1982-341621	19820122
JP 58000	986	A2	19830106	JP	1982-77396	19820507
JP 03016	358	B4	19910305			
US 45858	60	Α	19860429	US	1983-493051	19830509
US 45593	34	Α	19851217	US	1983-543880	19831020
US 49046	52	Α.	19900227	US	1985-785048	19851009
US 47314	43	Α	19880315	US	1986-889189	19860724
SU 15089	62	A3	19890915	SU	1987-4202592	19870519
JP 01308	286	A2	19891212	JP	1989-108256	19890427
JP 02111	751	A2	19900424	JP	1989-108255	19890427

US 4960889		Α	19901002	US	1990-462347		19900103
US 5026695		Α	19910625	· US	1990-461340		19900105
US 5110921		Α	19920505	US	1990-583304		19900917
US <sup>-</sup> 5594132		Α	19970114	US	1991-684194		19910412
US 5252731		Α	19931012	US	1992-831504		19920205
PRIORITY APPLN.	INFO.:			GB	1979-39985	Α	19791119
				GB	1980-4335	Α	19800208
				GB	1980-12991	Α	19800421
				GB	1980-22920	Α	19800714
				US	1980-205334	A2	19801110
				US	1980-206831	A3	19801114
				EP	1984-100915	Α	19801115
			•	EP	1987-104893	Α	19801115
				US	1981-261618	A2	19810507
				US	1982-341621	A3	19820122
				US	1982-428970	A2	19820930
•	ė			US	1983-489236	B1	19830428
				GB	1983-23034	Α	19830826
				US	1984-653041	Α3	19840921
				US	1985-785048	A3	19851009
				US	1986-889189	B3	19860724
				US	1987-127929	B1	19871202
				US	1990-462347	A3	19900103
					1990-461340	A3	19900105
				US	1990-583304	A3	19900917
GI			•	•	- 25		

$$\begin{array}{c|c} \text{RCH}_2\text{CXCCONH} & \text{S} \\ \parallel & \text{NOR}^1 & \text{CH} = \text{CH}_2 \\ & \text{CO}_2\text{R}^1 & \text{I} \end{array}$$

$$H_{2N}$$
 $S$ 
 $CCONH$ 
 $S$ 
 $CH = CH_{2}$ 
 $CO_{2}H$ 
 $CO_{2}H$ 
 $CO_{2}H$ 

AB Cephalosporins I [X = CO, protected CO; R = halogen; R1 = H, cycloalkenyl, (un) substituted alkenyl, alkyl, heterocyclic; R2 = H, protective group] were prepared Thus, benzhydryl 7-amino-3-vinyl-3-cephem-4-carboxylate.HCl was prepared from cephalosporin C in 6 steps and was acylated with BrCH2COC(:NOMe)CO2H to give I (R = Br, R1 = Me, R2 = CHPh2, X = CO) which was cyclized with thiourea and hydrolyzed to give the thiazolylacetamidocephem II. II had a min. inhibitory concentration against Proteus mirabilis of 0.05 μg/mL.

TT 79350-11-1P 79350-44-0P 79350-82-6P 86027-36-3P 90467-43-9P 90467-53-1P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and esterification of)

RN 79350-11-1 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,
7-[[(2-amino-4-thiazolyl) (methoxyimino)acetyl]amino]-3-ethenyl-8-oxo-,
monohydrochloride, [6R-[6α,7β(Z)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

● HCl

RN 79350-44-0 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2Z)-(2-amino-4-thiazolyl)(methoxyimino)acetyl]amino]-3-ethenyl-8-oxo-, monosodium salt, (6R,7R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

Na

RN 79350-82-6 CAPLUS
CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,
7-[[(2-amino-4-thiazolyl)[(carboxymethoxy)imino]acetyl]amino]-3-ethenyl-8oxo-, disodium salt, [6R-[6α,7β(Z)]]- (9CI) (CA INDEX NAME)

## Page 113

Absolute stereochemistry.

Double bond geometry as shown.

## ●2 Na

RN 86027-36-3 CAPLUS CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2-amino-4-thiazolyl)[[2-(diphenylmethoxy)-2-oxoethoxy]imino]acetyl]amino]-3-ethenyl-8-oxo-, monosodium salt, [6R-[6 $\alpha$ ,7 $\beta$ (Z)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

Na

RN 90467-43-9 CAPLUS CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2-amino-4-thiazolyl)[(2-propynyloxy)imino]acetyl]amino]-3-ethenyl-8-oxo-, monosodium salt, [6R-[6 $\alpha$ ,7 $\beta$ (Z)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Na

RN 90467-53-1 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2-amino-4-thiazolyl)(ethoxyimino)acetyl]amino]-3-ethenyl-8-oxo-, monosodium salt, [6R-[6 $\alpha$ ,7 $\beta$ (Z)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

Na

Absolute stereochemistry. Double bond geometry as shown.

HC1

RN

90467-54-2 CAPLUS
Pyridinium, 2-[[[[1-(2-amino-4-thiazolyl)-2-[(2-carboxy-3-ethenyl-8-oxo-5-CNthia-1-azabicyclo[4.2.0]oct-2-en-7-yl)amino]-2oxoethylidene]amino]oxy]methyl]-1-methyl-, chloride, [6R- $[6\alpha, 7\beta(Z)]$  - (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

Cl-

RN

90467-55-3 CAPLUS
Pyridinium, 3-[[[[1-(2-amino-4-thiazolyl)-2-[(2-carboxy-3-ethenyl-8-oxo-5-CN thia-1-azabicyclo[4.2.0]oct-2-en-7-yl)amino]-2oxoethylidene]amino]oxy]methyl]-1-methyl-, chloride, [6R- $[6\alpha, 7\beta(Z)]$  - (9CI) (CA INDEX NAME)

## Page 116

Absolute stereochemistry. Double bond geometry as shown.

● cl -

ANSWER 50 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN L9

ACCESSION NUMBER: 1984:68077 CAPLUS

DOCUMENT NUMBER: 100:68077

7-Acylamino-3-vinylcephalosporanic acid derivatives TITLE:

Fujisawa Pharmaceutical Co., Ltd., Japan PATENT ASSIGNEE(S):

Jpn. Kokai Tokkyo Koho, 20 pp. SOURCE:

CODEN: JKXXAF

DOCUMENT TYPE: Patent

LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE ---------\_\_\_\_\_ \_\_\_\_\_\_ \_\_\_\_\_ JP 58135894 Α2 19830812 JP 1983-9235 19830121 JP 05001271 **B4** 19930107 US 4487927 Α 19841211 US 1982-341621 19820122 PRIORITY APPLN. INFO.: US 1982-341621 Α 19820122 GB 1979-39985 Α 19791119 GB 1980-4335 A 19800208 GB 1980-12991 A 19800421 GB 1980-22920 A 19800714 US 1980-205334 A2 19801110 US 1981-261618 A2 19810507

Prepared by: Mary Hale @2-2507 Rem Bldg 1D86

GI

RCCONH 
$$C1CH_2COCCONH$$
  $S$   $CH=CH_2$   $CO_2H$   $CH=CH_2$   $CO_2H$   $CO_2H$ 

Nine cephalosporanic acid derivs. (I; R = aminothiazolyl; R1 = AB carboxyalkyl, protected carboxyalkyl; R2 = HO2C, protected HO2C) as the syn isomers were prepared I were effective bactericides at 50-2000 mg/day. Thus, 0.683 g (H2N)2CS and 1.84 g NaOAc were added to a suspension of 2.0 g syn-II in H2O at  $40^{\circ}$  and stirred 1.5 h to give 1.9 g syn-I (R = 2-aminothiazol-4-yl, R1 = MeO2CCH2; R2 = HO2C).

IT 88621-04-9P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

88621-04-9 CAPLUS RN

5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, CN 7-[[(2-amino-4-thiazolyl)[[2-[(4-nitrophenyl)methoxy]-2oxoethoxy]imino]acetyl]amino]-3-ethenyl-8-oxo-, monohydrochloride,  $[6R-[6\alpha,7\beta(Z)]]-(9CI)$  (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

● HCl

CAPLUS COPYRIGHT 2006 ACS on STN ANSWER 51 OF 52

1983:405441 CAPLUS ACCESSION NUMBER:

99:5441 DOCUMENT NUMBER:

7-Acylamino-3-vinylcephalosporanic acid derivatives TITLE:

Fujisawa Pharmaceutical Co., Ltd., Japan PATENT ASSIGNEE(S):

Jpn. Kokai Tokkyo Koho, 35 pp. SOURCE:

CODEN: JKXXAF

DOCUMENT TYPE: Patent Japanese LANGUAGE:

FAMILY ACC. NUM. COUNT: 9
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 58000986	A2	19830106	JP 1982-77396	19820507
JP 03016358	B4	19910305		
US 4423213	Α	19831227	US 1981-261618	19810507
PRIORITY APPLN. INFO.:			US 1981-261618 A	19810507
			GB 1979-39985 A	19791119
			GB 1980-4335 A	19800208
			GB 1980-12991 A	19800421
			GB 1980-22920 A	19800714
			US 1980-205334 A2	19801110
GI				

$$\begin{array}{c|c} R-CCONH & S & \\ \parallel & \parallel & \\ OR^1 & N & CH=CH_2 \\ \hline & R^2 & I \end{array}$$

AB Twenty title acids and salts (syn-I; R = aminothiazolyl with optional protecting group; R1 = carboxyalkyl, protected carboxyalkyl; R2 = carboxy, protected carboxy) were prepared I were effective bactericides at 50-2000 mg/day. Thus, 5 g syn-II was added to a suspension of POCl3 and DMF in THF under cooling, followed by 4.89 g III·HCl, and 9.2 g AcNHSiMe3 in EtOAc at -20° to -10° to give 3.7 g syn-I (R = 2-formamidothiazol-4-yl, R1 = Ph2CHO2CCH2, R2 = Ph2CHO2C).

IT 86027-36-3P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation and bactericidal activity of)

RN 86027-36-3 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,
7-[[(2-amino-4-thiazolyl)[[2-(diphenylmethoxy)-2-]
oxoethoxy]imino]acetyl]amino]-3-ethenyl-8-oxo-, monosodium salt,
[6R-[6α,7β(Z)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 52 OF 52 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1982:181061 CAPLUS

DOCUMENT NUMBER:

96:181061

TITLE:

7-Acylamino-3-vinylcephalosporanic acid derivatives,

pharmaceutical compositions containing them and their

starting compounds

INVENTOR(S):

Takaya, Takao; Takasugi, Hisashi; Masugi, Takashi;

Yamanaka, Hideaki; Kawabata, Kohji

PATENT ASSIGNEE(S):

Fujisawa Pharmaceutical Co., Ltd., Japan Eur. Pat. Appl., 285 pp.

SOURCE:

CODEN: EPXXDW

DOCUMENT TYPE:

LANGUAGE:

Patent English

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 30630 A2		19810624	EP 1980-107075	19801115
R: AT, BE, CH, DE,	FR, GB		NL, SE	
PRIORITY APPLN. INFO.:			GB 1979-39985 GB 1980-4335	19791119 19800208
			GB 1980-12991	19800421
	•		GB 1980-22920	19800714

GΙ

RXCONH 
$$O$$
  $CH = CH_2$   $CO_2R^1$ 

AB Vinylcephems I (R = optionally aminoheterocyclic, R2SO2NHC6H4; R 1 = H, protective group; R2 = alkyl; X = optionally substituted alkylene) were prepared Thus, I (R = 3-MeSO2NHC6H4, R1 = H, X = H2NCH, II) was obtained by acylating aminocephem with 3-MeSO2NHC6H4CH(NH2)CO2H. 7-Amino-3-vinyl-3-cephem-4-carboxylic acid was obtained from the hydroxymethylcephem via the chloromethyl derivative and the triphenylphosphonium iodide which was treated with CH2O. II had the min. inhibitory concentration against Staphylococcus aureus 209 P JC-1 of 1.56 μg/mL.

IT 79350-11-1P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation and bactericidal activity of)

RN 79350-11-1 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2-amino-4-thiazolyl) (methoxyimino)acetyl]amino]-3-ethenyl-8-oxo-, monohydrochloride, [6R-[6α,7β(Z)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

HCl

IT 79350-44-0P 79350-82-6P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and esterification of)

RN 79350-44-0 CAPLUS

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2Z)-(2-amino-4-thiazolyl)(methoxyimino)acetyl]amino]-3-ethenyl-8-oxo-, monosodium salt, (6R,7R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

Na

RN 79350-82-6 CAPLUS CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7-[[(2-amino-4-thiazolyl)[(carboxymethoxy)imino]acetyl]amino]-3-ethenyl-8-oxo-, disodium salt, [6R-[6 $\alpha$ ,7 $\beta$ (Z)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

●2 Na

Absolute stereochemistry.

● HCl

RN 90467-54-2 CAPLUS
CN Pyridinium, 2-[[[[1-(2-amino-4-thiazoly1)-2-[(2-carboxy-3-etheny1-8-oxo-5-thia-1-azabicyclo[4.2.0]oct-2-en-7-yl)amino]-2 oxoethylidene]amino]oxy]methyl]-1-methyl-, chloride, [6R-[6α,7β(Z)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

Ocl -

Absolute stereochemistry.

● Cl -

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FILE 'REGISTRY' ENTERED AT 14:28:31 ON 03 MAR 2006 E CEFDINIR/CN 5

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L2 STR 91832-40-5

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L4 STR L3

L5 7 S L2 OR L3 OR L4

L6 166 S L2 OR L3 OR L4 FUL

L7 SCR 2127

L8 55 SEARCH L7 SUB=L6 FUL

FILE 'CAPLUS' ENTERED AT 14:32:44 ON 03 MAR 2006 L9 52 S L8

FILE 'CAOLD' ENTERED AT 14:34:02 ON 03 MAR 2006 L10 0 S L9

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L2 STR

NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

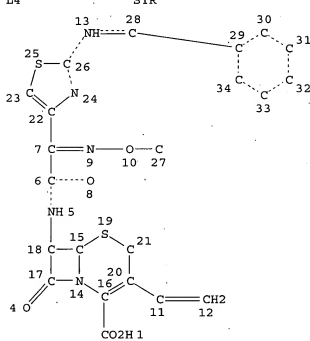
GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 25

STEREO ATTRIBUTES: NONE L3 STR

NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 27

STEREO ATTRIBUTES: NONE L4 STR



NODE ATTRIBUTES:

DEFAULT MLEVEL IS ATOM

DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED

NUMBER OF NODES IS 32

STEREO ATTRIBUTES: NONE

L6 166 SEA FILE=REGISTRY SSS FUL L2 OR L3 OR L4

L7 SCR 2127

L8 55 SEA FILE=REGISTRY SUB=L6 SSS FUL L7

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